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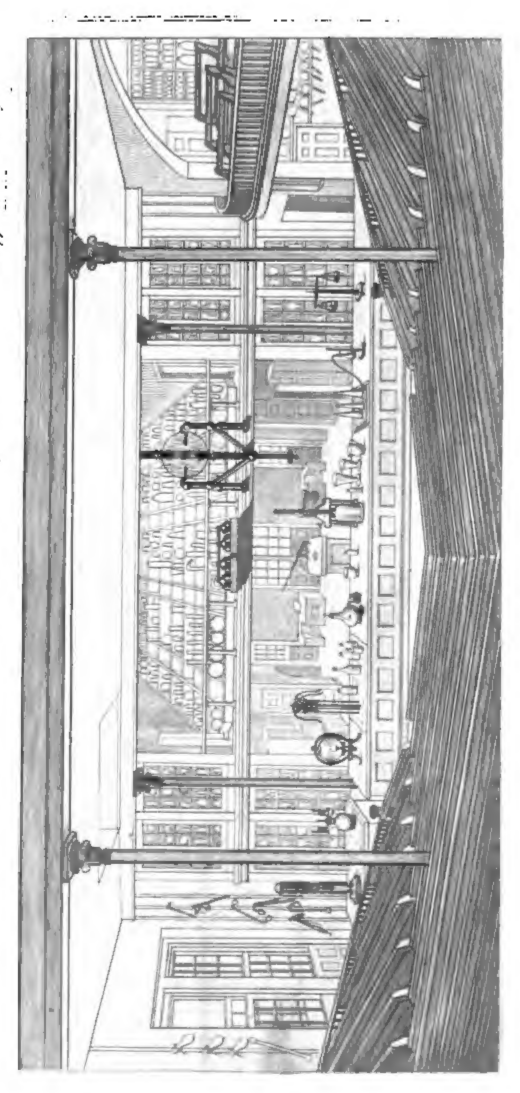
The
Charles Motley Clark
Memorial











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COMPENDIUM

OF THE

COURSE OF CHEMICAL INSTRUCTION,

IN THE

MEDICAL DEPARTMENT

OF THE

UNIVERSITY OF PENNSYLVANIA.

ROBERT HARE, M.D.

IN TWO PARTS.

PART I.

COMPRISING THE

CHEMISTRY OF HEAT AND LIGHT, AND THAT OF INORGANIC SUBSTANCES, USUALLY CALLED INORGANIC CHEMISTRY.

FOURTH EDITION.

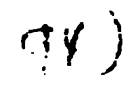
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PREFACE

TO THE FIRST EDITION.

WHERE a subject cannot be followed by a reader without study, it would seem unreasonable to expect that, without some assistance, it should be followed at a lecture. Under this impression, from the time that I became a lecturer, I applied myself so to improve and multiply the means and methods of experimental illustration, as to render manipulation easier, and the result more interesting and instructive.

But notwithstanding all my efforts, there remained obstacles to be surmounted. However striking might be the experimental illustration of a property or principle, the rationale might be incomprehensible to a majority of my class, unless an opportunity for studying it were afforded them.

Again, some of my contrivances, which greatly facilitated my experiments, were too complex to be understood without a minuteness of explanation, which, even if it were useful and agreeable to some of my hearers, much be useless and irksome to others; and to such minutiæ I have not deemed it expedient to exact attention.

A chemical class, in a medical school, usually consists of individuals, who in the widely with respect to their taste for chemistry, and in opinion as to the extent to which it may be practicable or expedient for them to mean it. There is also much disparity in the opportunities which they may have enjoyed, of acquiring some knowledge of this science, and of others which are subsidiary to its explanation. Hence a lecturer may expatiate too much for one portion of his auditors, and yet be too concise for another portion. While to the adept he may often appear trite, to the novice he may often appear abstruse.

Some pupils, actuated by a laudable curiosity, under circumstances permering its indulgence, may desire an accurate knowledge of the apparatus to which my experimental illustrations are facilitated: other pupils may feel themselves justified, perhaps necessitated, not to occupy their time with the acquisition of any knowledge which is not indispensable to graduation.

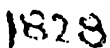
After some y s' experience of the difficulties abovementioned, I came

that the time spent in the lecture room might be rendered ple, if students could be previously apprized of the chain

of ideas, or the apparatus and experiments, to be subjected to attention at each lecture; especially as the memory might afterwards be refreshed by the same means. In consequence of this conviction, the minutes of my course of instruction were printed; and subsequently a work, comprising engravings and descriptions of the larger portion of such of my apparatus and experiments, as could in this way be advantageously elucidated. Encouraged by the success of my plan, I am now preparing an edition which will be still more extensive. The work thus expanded, I have entitled "A Compendium of the Course of Chemical Instruction in the Medical School," &c.

There will be much matter in the Compendium, respecting which I shall not question candidates at the examination for degrees. With the essence of the larger part, I shall undoubtedly expect them to be acquainted; but other portions have been introduced, that I may not be obliged to dwell upon them in my lectures, and that attention to them may be optional on the part of the students. To designate the portion of the work, respecting which candidates for degrees will not be questioned, I have had it printed in a smaller type, excepting where it was too much blended with subjects of primary importance to be separated. I wish it, however, to be understood, that I shall expect attention to the parts thus distinguished, so far as they may be necessary to a comprehension of the rest. Thus, although I do not deem it to be a part of my duty to question a pupil on pneumatics, I shall expect him to understand the influence of atmospheric pressure upon chemical reaction, and in pneumato-chemical operations.

One great and almost self-evident advantage, resulting from my undertaking, I have yet to mention; I allude to the instruction which students may derive from the Compendium, either before or subsequently to their attendance on my lectures, and especially during the period which intervenes between their first and second course.



PREFACE

TO THE FOURTH EDITION.

Tex suggestions, which were made in the Preface to the first edition of the Compendium, respecting the necessity of an appropriate text book, to ad and extend the instruction afforded by the course of chemical lectures, deinered in the Medical Department of the University of Pennsylvania, have anguired additional force since that Preface was written. During the twelve intervening years the boundaries of those portions of human knowledge over which Chemistry has established a rightful domain, have undergone an extrasion commensurate with the time. It is, of course, proportionably more difficult to do justice to the whole of the wonderful region comprised within thuse boundaries in sixty lectures delivered within four months. the attention of the student was alternately claimed by six professors; but laterly, the claims of a seventh professor have been added to those prevery established. Nevertheless, I am under the impression, that with the assistance which my text books are competent to afford, my course of lectirs, brief as it is, may be more serviceable to a student who makes due we of those text books, than it could prove, were its duration doubled, withcut bing associated with treatises made expressly for the purpose of amplithe information partially afforded by my lectures, or of remedying ver acvitable omissions.

Have a been prevented by indisposition from commencing this work as early as expedent. I am under the necessity of issuing that part which release to Caloric, Light, and Inorganic Chemistry first. Dynamic Electricity, comparing Galvanism or Voltaic Electricity, and Electro-magnetism, tax no issue already issued, I shall in the next place republish my Treatise on Mechanical Electricity. Then to complete the new edition of my text terms only Organic Chemistry will remain to be reprinted. On this branch I tage to turnish a treatise before I reach that part of my course of lectures, in which it becomes the object of attention.

I am in hope's that numbering the paragraphs, an excellent expedient remed to by me for the first time in this edition, will be found advantageous to the reader, by rendering references from one part of the work to another inconvenient, and consequently more frequent.



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DEFINITIONS OF CHEMISTRY.

It is natural that a person whose attention may be directed to chemshould inquire of what does it treat, or how is it to be defined or a guished from other sciences?

Agreeably to the definition given in the second page of the Compen chemistry treates of those phenomena and operations of nature which from reaction between particles of inorganic matter.

I subjoin several other definitions from some of the most celel modern writers on chemistry.

Thomson defines chemistry to be "the science which treats of events or changes in natural bodies, which are not accompanied by sible motions."

Henry conceives that "it may be defined, the science which invest the composition of material substances, and the permanent change constitution, which their mutual actions produce."

According to Murray, "it is the science which investigates the c nations of matter, and the laws of those general forces, by which combinations are established and subverted."

Brande alleges "that it is the object of chemistry to investiga changes in the constitution of matter, whether effected by heat, mixtu other means."

According to *Ure*, "chemistry may be defined that science, the of which is to discover and explain the changes of composition that among the integrant and constituent parts of different bodies."

The definition given by *Berzelius* is as follows:—"Chemistry science which makes known the composition of bodies, and the mans which they comport with each other."

COMPENDIUM

O.

CHEMICAL INSTRUCTION,

&c.

INTRODUCTION.

1. The phenomena and operations of the material world spear to be dependent on certain properties in the partides or masses of matter which enable them to exercise • reciprocal influence. Without this reciprocal action, which I would prefer to call reaction,* every particle or was would be as if no other existed, and could itself have meficient existence.

2. The reciprocal action or reaction, thus inferred to exist may be distinguished as taking place between masses, between a mass and particles, and between particles only.

3. Reaction between massest is sublimely exemplified in the solar system, by that attraction between the sun and planets, by which they are made to revolve in their orbits.

4. Reaction between a mass and particles is exemplified by the reflection, refraction, and polarization of light.

5. Reaction between particles is exemplified by a fire, or the explosion of gunpowder.

Definition of Natural Philosophy, Chemistry, and Physiology.

6. Natural Philosophy, in its most extensive sense, treats of physical reaction generally. In its more limited and

La Mochanics, action is said to produce reaction; but in the case of an innate which mutually causes different portions of matter to be self attractive, Type word mass, I mean a congeries of particles capable of producing some

substively, to which severally they would be incompetent.

usual acceptation, it treats of those phenomena and operations of nature, which arise from reaction between masses, or between a mass and particles.

7. Chemistry treats of the phenomena and operations of an nature, which arise from the reaction between the particles

of inorganic matter.

8. Physiology treats of the phenomena and operations, which arise from the reaction of the masses or atoms of organic or living bodies.

OF CHEMICAL REACTION.

ť

9. Reaction between particles, or chemical reaction, is distinguished into repulsive reaction or repulsion, and attractive reaction or attraction.

OF REPULSIVE REACTION OR REPULSION.

A Priori Proofs that there must be a Matter in which Repulsion exists as an Inherent Property.

- 10. Matter may be defined to be that which has properties. We know nothing of matter directly. It is only with its properties that we have a direct acquaintance. It is from our perception of matter, through the powers or properties by which it affects our senses, that we believe in its existence.
- dent as that of the matter which, in obedience to their successive predominancy, may be seen either to cohere, in solids, with great tenacity, or to fly apart with explosive violence in the state of a vapour. The existence of repulsion and attraction being proved, it must be admitted that they are properties of matter; since the existence of a property, independently of matter, is inconceivable. But being of a nature to counteract each other, the repellent and attractive powers cannot coexist in particles of the same kind, and consequently must belong to particles of different kinds. There must, therefore, be a matter endowed with repulsion, distinct from that which is endowed with attraction.
- 12. I conceive that the phenomena of chemistry demonstrate that there are at least the three following properties, which, from their obvious incompatibility, cannot belong to the same elementary particles.

13. 1st. An innate property of reciprocal attraction.

14. 2d. An innate property of counteracting attraction

directly, by imparting reciprocal repulsion.

15. 3d. An innate property of imparting an attraction, variable in its force, and limited and contingent in its duration.

16. I presume that there must be at least three different kinds of matter, to each of which, one of the properties thus specified innately appertains.

17. The permanent and unvarying attractive power is exemplified by gravitation, and, as modified by circum-

stances, by tenacity, or cohesion.

18. It resides, undoubtedly, in every kind of matter endowed with weight, and consequently in all that is considered as material by the mass of mankind.

19. It must likewise act between each of those imponderable principles which I am about to mention, and all

other matter, whether ponderable or imponderable.

- 20. The power of imparting reciprocal repulsion to penderable matter is supposed by chemists generally to belong to certain imponderable material reciprocally repulsive particles, constituting the cause of heat, called caloric.
- 21. The power of indirectly counteracting attraction, and substituting for it a contingent and variable attraction, appears to belong to electricity. Light also appears to exercise a modifying influence.
- 22. Thus we have reason to infer the existence of at least three imponderable substances—electricity, caloric, and light—each consisting of particles reciprocally repulsive. yet attractive of other matter, and probably more or less attractive of each other.

OF CALORIC.

Experimental Proofs of the Existence of a material Cause of Calorific Repulsion.

23. It has been ascertained that ice melts and water bees at the temperature of 32° of Fahrenheit's thermomer. If at this temperature, which is called the freezing ice in a divided state, as in that of snow for interpretation, be mingled with an equal weight of water at 172°,

the ice will be melted, and the resulting temperature will be 32°; but if equal weights of water be mingled at those temperatures, the mixture will have the mean heat of 102°.

24. It follows that a portion of heat becomes latent in the aqueous particles during the liquefaction of the ice, sufficient to raise an equal weight of water one hundred and forty degrees. In this case the ice is supposed to combine with material calorific particles, innately endowed with a power of reciprocal repulsion, and likewise with that of combining with ponderable matter. Hence water is considered as a combination of ponderable particles, endowed with a reciprocally attractive power, and imponderable particles endowed with a reciprocally repellent power; so that, in obedience to the power last mentioned, the compound atoms, instead of cohering as in the solid state, move freely among each other, forming consequently a liquid.

25. In all cases of liquefaction or fusion which have been examined, analogous results have been observed; whence it is generally believed that whenever a solid is converted into a liquid, its particles unite with a portion of the material cause of heat, which becomes latent, as in the case of ice in melting. The evidence is equally strong in favour of the inference that in passing from the liquid to the aëriform state, ponderable matter combines with, and renders latent even a larger quantity of heat in proportion

to its weight, than in cases of liquefaction.

26. When, by means of a thermometer, we observe the rise of temperature in water exposed to a regular heat, as when placed in a cup upon a stove, we find that nearly equal increments of heat are acquired in equal times, until the boiling point is attained. Subsequently, the cup being open so as to allow the steam to escape freely, no further rise of temperature will be found to ensue; but in lieu of it, steam will be evolved more or less copiously, in proportion to the activity of the fire. Since from the time the water boils it ceases to grow hotter, it may be fairly presumed that the steam generated during the ebullition, although of a temperature no higher than 212°, contains, in a latent state, the caloric which meanwhile enters the liquid. This presumption is fully justified by the fact, that if any given weight of steam be received in a quantity of

cool water ten times heavier, it will cause in it a rise of

temerature of nearly one hundred degrees.

In the heat which would raise ten parts of water to lodegrees, would, if concentrated into one of those parts, rise it to 1000 degrees nearly, which is about equal to a red heat. It follows, therefore, that as much heat is abouted in producing steam, as would render the water of which it consists red-hot, if prevented from assuming the utilorn state.

28. These facts and deductions induce chemists generally to believe that the cause of calorific repulsions is material; that it consists of a fluid, of which the particles are refrepellent, while they attract other matter; that by the mion of this fluid with other matter, a repulsive property is imparted, which counteracts cohesion, so as to cause, successively, expansion, fusion, and the aëriform state; and further, that it is by the afflux of the calorific matter that the sensation of heat is produced, while that of cold results from its efflux.

Acceptation of the term Caloric.

29. If we place a small heap of fulminating mercury upon the face of a hammer, and strike it duly with another hammer, an explosion will ensue so violent as to cause a visible indentation in the steel surface. This explosion, agreeably to the premises, can only be explained by suppreing the evolution of a great quantity of the material Were an equal quantity of red-hot sand to b: suddenly quenched with water, the effect would be comparatively seeble. We may, therefore, infer that the fulmnating powder, though cold, contains more of the cause of heat than a like quantity of red-hot sand. Hence it would follow from using the word heat in the sense both of cause and effect, that there is more heat in a cold body than in a hot one, which in language is a contradiction. On this account it was considered proper by the chemists of the Lavoisierian school, to use a new word, caloric, to designate the material cause of calorific repulsion.

Experimental Illustration.

3. A portion of fulminating mercury exploded between two hammers.

ORDER PURSUED IN TREATING OF CALORIC.

Expansion.—Modification of the effects of Caloric by Atmospheric Pressure.—Capacities for Heat, or Specific Heat.— & Slow Communication of Heat, comprising the Conducting Process and Circulation.—Quick Communication of Heat, or Radiation.—Means of producing Heat, or rendering Caloric sensible.—Means of producing Cold, or rendering Caloric laters. States in which Caloric exists in Nature.

EXPANSION.

OF THE EXPANSION OF SOLIDS, LIQUIDS, AND ELASTIC FLUIDS, AND ON THE OPPONENT AGENCY OF ATMOSPHERIC AND OTHER PRESSURE.

Expansion of Solids.

31. A ring and plug, which when cold fit each other, cease to do so when either is heated; and a tire when red-hot is made to embrace a wheel otherwise too large for it.

Pyrometer, in which the Extension, in length, of a Metallic Bar is rendered sensible by a Combination of Levers.

32. The influence of temperature on the length of a metallic wire may be rendered evident by means of the instrument, of which fig. 1, in the oppo-

site engraving is a representation.

33. WW, represents a wire, beneath which is a spirit lamp consisting of a long, narrow, triangular vessel of sheet copper, open along the upper angle, so as to receive and support a strip of thick cotton cloth, or a succession of wicks. By the action of the screw at S the wire is tightened, and by its influence on the levers, the index I is raised. The spirit lamp is then lighted and the wire enveloped with flame. It is of course heated and expanded, and, allowing more liberty to the levers, the index upheld by them falls.

34. By the action of the screw the wire may be again tightened, and, the application of the lamp being continued, will again, by a further expansion, cause the depression of the index; so that the experiment may be

repeated several times in succession.

35. Since this figure was drawn, I have substituted for the alcohol lamp the more manageable flame of hydrogen gas, emitted from a row of apertures in a pipe supplied by an apparatus for the generation of that gas. See fig. 2.

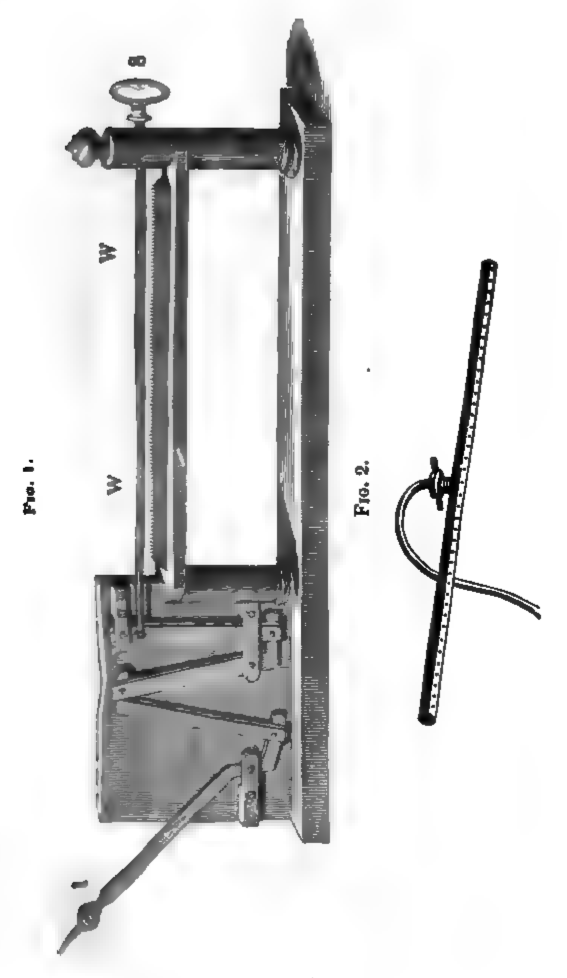
36. If, while the index is depressed by the expansion, ice or cold water be applied to the wire, a contraction immediately follows so as to raise the

index to its original position.

37. Metals are the most expansible solids, but some are more expansible than others.

38. The following table, abstracted by Turner from that furnished by Lavoisier, will show the increase of bulk obtained by glass and various metals in rising in temperature from 32° to 212°.

Instrument
demonstrating the Power of Caloric in expanding a Metallic Rod.



(Pege 6.)



Elongation when keated man of Substances. from 32° to 212°. subs without lead, mean of three specimens 1-1115 of its length. in first glass, 1-1248 1.581 L mean of two specimens, 1.532rea, forged, 1-819 1.812 spered steel, 1-927 med steel, 1.807 fladia, 1-516 f Falmouth, 1-462 , mean of three specimens, 1-602 num, determined by Borda, 1.1167

A Pyrometers have been made of platinum, in one of which, invented Desical, changes in the length of a cylinder of this metal, arising from wature, are made sensible by the motion of a lever associated with it, which acts as an index. In the other, a bulb is formed of platinum, be degree of heat is inferred from the quantity of air expelled.

L The use of this air pyrometer is burdened by the necessity of meament and calculation to ascertain the result. This might be very much med by the use of a sliding rod and air-gauge. The retraction of ed might be made to compensate the expulsion of air, while divisions mde on it would indicate the quantity.

sperimental Illustration of the different Expansibility of Metals.

11. That the expansibility of one metal may exceed that another, may be rendered apparent by soldering tober, face to face, two thin strips, one iron the other On exposure to heat, the compound strip, thus estituted, assumes the shape of an arch. The brass, buch is the more expansible metal, forms the outer and of urse larger curve.

pposed Exception to the Law that Solids expand by Heat in the case of Clay, which contracts in the Fire.

12. The phenomena do not justify us in considering the traction of clay from heat as an exception to the geal law. In the first instance clay shrinks by losing ter, of which the last portions are difficult to expel. exect place a chemical union takes place between the cipal ingredients, silica and alumina, which is rendered examplete in proportion to the duration and intensity fire. It may be presumed that the vitreous compound, which would result from a complete fusion as combination of the constituents, would be as expansible other vitreous substances.

Experimental Illustration.

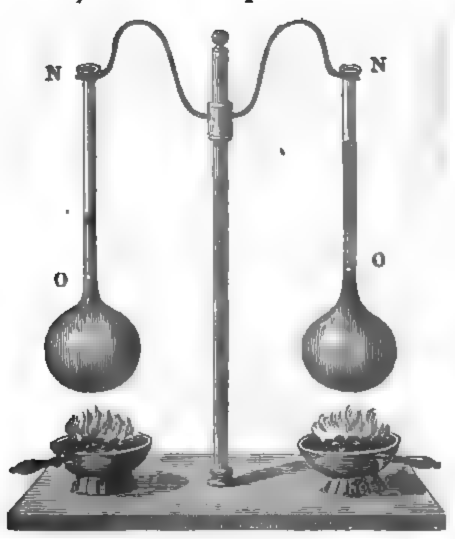
43. The contraction produced by heat in cylinders clay shown by means of the ingenious but inaccurate pyr meter of Wedgwood.

Expansion of Liquids or non-elastic Fluids.

44. The word fluid applies to every mass that will flodistribute itself equally in obedience to its own weight self-repulsion.

45. Ponderable fluids are either elastic or non-elastic Latterly the term liquid has been employed to designathose fluids which are, like water, alcohol, and oil, devo of elasticity, a property which, in due time, I shall definand illustrate.

Liquids are expanded when their Temperature is raised, as some Liquids are more expansible than others.



Medical of the same shape and dimensions as represented in the preceding figure. Let one of them, that on the left for instance, be supplied with as much alcohol as will occupy it to the level designated by the letters O.O. Let the vessel on the right be occupied with water to the same level, the height of the liquid in each being made to correspond with a little fillet of white paper secured about the neck. Inder each vessel, place equal quantities of charcoal, burning with a similar degree of intensity; or preferably, surround the bulbs simultaneously with hot water in an oblong vessel of suitable dimensions. The liquids in each vessel will be expanded so as to rise into the necks; but the alcohol will rise to a greater height than the water.

17. The dilatation of the following liquids, by a change of temperature from 32° to 212°, is as follows—alcohol 1-9, and a change acid 1-9, fixed oils 1-12, sulphuric ether 1-14, oil of temperature 1-14, sulphuric or muriatic acid 1-17, brine

1-23 nearly, mercury about 1-55.

48. The rate of expansion for liquids increases with the temperature; as if their particles, by becoming more remote, lost some of their ability to counteract the repulsive valuence of caloric.

49. The number associated with each of the substances in the following list, shows its melting point as estimated by Fahrenheit's scale. One degree of Daniell's pyrometer, (39) by which the temperatures above 600° were measured, is

calculated to be equal to seven of Fahrenheit.

In. Cast iron 3479°, gold 2590°, silver 2233°, brass 1869°, anumony \$10°, zinc 648°, lead 606°, bismuth 497°, tin 442°, sulphur 218°, beeswax 142°, spermaceti 112°, phosphorus 118°, tallow 92°, olive oil 36°, milk 30°, blood 25°, water 271°, oil of turpentine 14°, mercury—39°, nitric acid—15!°, sulphuric ether—46°.

Exception to the Law that Liquids expand by Heat.

I. The bulk of water diminishes with the temperature, with it reaches 39° nearly. Below this point, it expands at grows colder, and in freezing increases in bulk one. This wonderful exception to the law that liquids expend by heat, appears to be a special provision of the Dery for the preservation of aquatic animals; for were

water to increase in density as it approaches the poin of congelation, the upper stratum would continue to sind as refrigerated in bodies of water below 39°, as well as it others. Hence a whole river, lake, or sea might, in high latitudes, be rendered too cold for animal life; and finally be so far converted into ice, as not to thaw during the ensuing summer. Subsequent winters co-operating, the whole might be consolidated so as never to thaw. in consequence of the peculiarity in question, the cold est stratum, in a body of water below 39°, remains at top until, if the cold be adequate, congelation ensues. buoyant sheet of ice, which results in this case, forms effec tively a species of winter clothing to the water beneat it; and, by augmenting with the frost, opposes an increas ing obstacle to the escape of caloric from the water which it covers.

Expansion of Aëriform Fluids.

52. Aëriform fluids are much more expansible than liquids. In order, however, to appreciate the changes of bulk which they may be observed to sustain, it is necessary to understand the influence which the pressure of the atmosphere has upon their density, independently of temperature. The simple influence of heat, in expanding them may be illustrated by holding a hot iron over the thermometer of Sanctorio, represented in the following figure.

Thermometers.

53. The invention of the thermometer is ascribed to Sanctorio. The principle of that form of the instrument which he contrived may be understood from the following article.

11

species of Air illustrated by the Air Thermometer of Sanctorio on a large Scale.



ed by a ring and an upright wire with its neck downwards, so as to have its orifice beneath the surface of the water in a small glass jar. A heated iron being held over the matrass, the contained air is so much increased in bulk, that, the vessel being inadequate to hold it, a partial escape from the orifice through the water ensues. On the removal of the hot iron, the residual air regains its previous temperature, and the portion expelled by the expansion is replaced by the water.

55. If in this case the quantity of air expelled he so regulated, that when the remaining portion returns to its previous temperature, the liquid rises about half way up the stem, or neck, the apparatus will constitute an air thermometer. For whenever the temperature of the external air changes, the air in the bulb of the matrass

distriction of bulk, and consequently, in a proportionate degree, influction of bulk, and consequently, in a proportionate degree, influction to the liquid in the neck. As elastic fluids are dilated abit, in proportion to the temperature, and are also much more expansion but has the temperature would be very accurate, as well as temperature. On this account, however, it was never of much as temperature. On this account, however, it was never of much as temperature. On this account, however, it was never of much account as a possible control of the instrument assumed a form now generally employed, the principle of which is explained. (45.) So letter following pages I shall give engravings and descriptions of a form of the thermometer used in the laboratory, of the self-registering theoreter, of the differential thermometer, and of an apparatus which through the difference between it and Sanctorio's thermometer.

57. Agreeably to the example of my predecessor and preceptor Dr. velouse. I have been accustomed to exhibit to my class the blowing and lag of a thermometer. Of this process an account is subjoined.

The tribes used in constructing thermometers are made at almost all the glass two having usually a capillary perforation. They are made by rapidly drawing to hollow glass globe while red-hot, by which means it is changed into a long inducal string of glass, in the axis of which a perforation exists, in consequence to eavily of the globe. When a thermometer tube is softened by exposure to a two extend by a blow-pipe, a bulb may be blown upon it. While the bulb is still the other end of the tube is immersed in mercury, or in spirit, according to the start and the substitute of the instrument is intended. As the bulb cools, the air within the liquid, and the allows the liquid to enter, in obedience to the pressure of the liquid. The bulb thus becomes partially supplied with the liquid, which is think in order to expel all the air from the cavity of the bulb and perforation.

The orifice being again depressed into the liquid, when the whole becomes cold the liquid will fill the cavity of the bulb. This result will be hereafter fully explained and illustrated. The open end of the tube being now heated, is drawn out into a filament with a capillary perforation. The bulb being raised to a temperature above the intended range of the thermometer, so as to expel all the superabundant liquid, the point is fused so as to seal the orifice hermetically, or in other words so as to be perfectly air-tight. In the next place, the bulb is to be exposed to freezing water, 🖨 and the point to which the liquid reaches in the capillary perforation marked. In like manner the boiling point is determined, by subjecting the bulb to boiling water. The distance between the freezing and boiling points, thus ascertained, may be divided according to the desired graduation.

59. The scale of Reaumur requires 80 divisions, that of Celsius 100, Fahrenheit's 🛌 180. The graduation of Celsius is the most rational; that of Fahrenheit the least so, although universally used in Great Britain and the United States. The degrees of these scales are to each other obviously, as 80, 100, and 180; or as 4, 5, and 9.

Hence it is easy to convert the one into the other by the rule of three.

60. It should, however, be observed that the scales of Celsius and Reaumur commence at the freezing of water, all above that being plus, all below it minus; while the scale of Fahrenheit commences at thirty-two degrees below freezing. Hence in order to associate correctly any temperature noted by his thermometer with theirs, we must ascertain the number of degrees which the mercury is above or below freezing, and convert this number into one equivalent to it by their gradustion; and conversely, after changing any number of degrees of theirs into his, we must consider the result as indicating the number of degrees above or below 32 on his scale.

61. The process above described for the construction of a thermometer, is equally applicable whether the bulb be filled with alcohol or mercury. Each of these liquids has peculiar advantages. Mercury expands most equably. Equal divisions on the scale of the mercurial thermometer will more nearly indicate equal increments or decrements of temperature. Mercury also affords a more extensive range; as it does not boil below 656°, nor freeze above—39°, of Fahrenheit's thermometries scale.

62. Alcohol, being more expansible than mercury, is more competent to detect slight changes. It boils at 176° of Fahrenheit, and for its congelation is alleged to require— 90° of the same scale. As this temperature is below any ever observed in nature, and can only be attained by an extremely difficult process, latterly discovered by Bussier, it can hardly ever happen that an alcoholic thermometer will not be found competent to measure any degree of cold which chemists have a motive for determining. Besides those above mentioned, a thermometric scale has been used in Russia, which bears the name of its author, Delisle. In this, zero is at the boiling point of water, and five of his graduations are equal to six of Fahrenheit's.

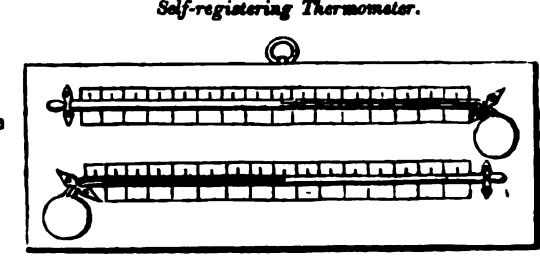
Laboratory Thermometer.

63. The thermometers used in laboratories, are usually constructed so as to have a portion of the wood or metal, which defends them from injury and receives the graduation, to move upon a hinge, as represented in the adjoining figure.

64. This enables the operator to plunge the bulb into fluids, without introducing the wood or metal, which would often be detrimental either to the process or to the instrument, if not to both.

65. The scale is kept straight by a little bolt on the back of it, when the thermometer is not in use.

Self-registering Thermometer.



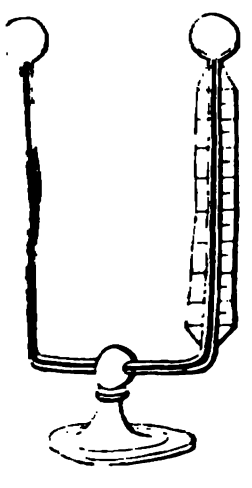
The figure represents a self-registering thermometer. It comprises necessarily a serious and a spirit thermometer, which differ from those ordinarily used, in having the stems horizontal and their bores round; also large enough to admit a cylinder of samel in the bore of the spirit thermometer, and a cylinder of steel in the bore of the mercural thermometer. Both the cylinder of enamel and that of steel must be meanly of the same diameter with the perforations in which they are respectively unated, as is consistent with their moving freely in obedience to gravity, or any patte impulse.

The order to prepare the instrument for use, it must be held in such a situation, what we ensured may subside as near to the end of the alcoholic column as possible, set all remaining within this liquid. The steel must be in contact with the mer-

ery, but not at all immersed in it.

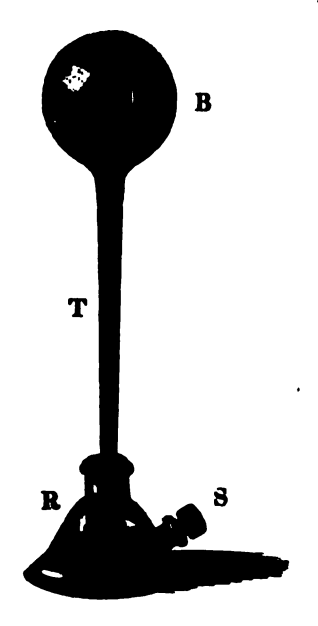
© On this account the bulbs of the thermometers are placed at opposite ends of the the upon which they are secured; so that when this plate is made to stand up on end. 12 such manner as to have the bulb of the mercurial thermometer lowerment, that of the spirit thermometer will be uppermost. Under these circumstances, mpelled by gravity, the steel cylinder will subside upon the surface of the mercurial ma while the cylinder of enamel will sink within the little column of spirit, which means it, till it reaches the surface of that column. The instrument being, wher the object is attained, suspended in a horizontal position, as represented in the from if in consequence of its expansion by heat, the mercury advance into the whe the same! moves before it; but should the mercury retire during the absence of to deriver, the steel does not retire with it. Hence, the maximum of temperature, a the interior, is discovered by noting the graduation opposite the end of the cylinar morest the mercury. The minimum of temperature is registered by the enamel, which retreets with the alcohol when it contracts, but, when it expands, does not since with it. The enamel must retire with the alcohol, since it lies at its marga, and cannot remain unmoved in the absence of any force competent to extricate a seem a liquid, towards which it exercises some attraction. But when an opposite movement takes place, which does not render its extrication from the liquid necesmy to in being stationary, the enamel does not accompany the alcohol. Hence the manner of temperature, which may have intervened during the absence of the observer, is discovered by ascertaining the degree opposite the end of the enamel nearest to the end of the column of alcohol.

Leslis's Differential Thermometer.



On This instrument consists of a glass tube nearly in the form of the letter U, with a bulb at each termination. In the bore of the tube there is some liquid, as, for instance, coloured sulphuric acid, alcohol, or ether. When such an instrument is exposed to any general alteration of temperature in the surrounding medium, as in the case of a change of weather, the air in both bulbs being equally affected, there is no movement produced in the fluid; but the opposite is true, when the slightest change of temperature exclusively affects one of the bulbs. Any small bodies situated at different places in the same apartment warmed by a fire, will show a diversity of temperature, when severally applied to the different bulbs.

Difference between an Air Thermometer and a Differential Thermometer, illu upon a large Scale.



70. The adjoining figure rept an instrument, which acts ac thermometer, when the stoppi removed from the tubulure in th cal recipient, R; because in the whenever the density of the phere varies either from chan, temperature, or barometric pro hereafter to be explained, the of the alteration will be indicate an increase or diminution of the occupied by the air in the bu and of course by a corresp movement of the liquid in the T. But when the stopple is in its the air cannot, within either ca the instrument, be affected by c in atmospheric pressure: no changes of temperature which (equally on both cavities, produ movement in the liquid which rates them. Hence, under the cumstances, the instrument is tent to act only as a differentia mometer.

MODIFICATION OF THE EFFECTS OF CALORIC BY AT PHERIC PRESSURE.

Digression to demonstrate the Nature and Extent of Atmospher Pressure.

Experimental Proof that Air has Weight.

71. The air being allowed to replent exhausted globe, while suspended from a beam and accurately counterpoised, can to preponderate.

72. By a temporary communication wi air pump, by means of a screw with whis furnished, a glass globe is exhausted of It is then suspended to one arm of a scale and accurately counterpoised. Being thu pared, if by opening the cock the air lowed to re-enter the globe, it will preparet; and if a quantity of water, adequates and if a quantity of water, adequates the equilibrium, be introduced small vessel, duly equipoised by a coweight applied to the other arm of the the inequality in bulk of equal weights and water will be satisfactorily exhibited.

Definition of Elasticity.

the power which bodies have to resume their shape, position, or the censation of constraint, is called clasticity. The degree in teny body possesses this power is not to be estimated by the force, the perfection of its recoil. A coach spring is far more powerful, not more clastic, than a watch spring.

. Elasticity is erroneously spoken of as a varying property in the air, in common with aëriform fluids in general, appears to be always

sly elastic.

. As a property distinguishing aëriform fluids from liquids, elasticity eye the idea of a power in a given weight of a fluid to expand or to not with the space in which it may be confined, producing at the same a pressure on the internal surface of the cavity, or any object within tensily as the space.

bistoms and Extest of the Pressure of the Atmosphere experimentally demonstrated.

PRELIMINARY PROPOSITION.

For the pressure of any fluid on any area assumed within it, the pressure of a coof any other fluid may be substituted, making it as much higher as lighter, as leaser as homewer; or in other words, the heights are inversely as the gravities.

Experimental Illustration in the case of Mercury and Water.



77. If into a tall glass jar, such as is represented in the adjoining figure, a glass cylinder, C, (like a large glass tube open at both ends) were introduced—on filling the jar with water, this liquid would of course rise in the cylinder to the same baight as in the der to the same height as in the jar; but, if, as in the figure, before introducing the water, the bottom of the jar be covered with a stratum of mercury, two meches deep, so as to be sufficiently above the open end of the cylinder, it must be evident that the water will be prevented from entering the cylinder by the interposition of a heavier liquid. But as the pressure of the water on the mercury outside of the cylinder is unbalanced by any pressure from water within the cylinder, the mercury within will rise, until, by its weight, the external pressure of the water is compensated. When this is effected, it will be seen, on comparing, by means of the scale, S, the height of the two liquids, that for every inch of elevation acquired by the mer-cury, the water has risen more than a foot, since the weight of mercury is to that of water, as 13.6 to 1.

78. It may be demonstrated that the pressure of the column of mercury is at equivalent to that of a column of water having the same base, and an all equal to that of the water in the jar, by filling the cylinder with water. I then be seen, that, when the water inside of the cylinder is on a level wit water on the outside, the mercury within the cylinder is also on a level wit

mercury without.

79. It is, therefore, obvious, that the elevation of the column of mercury, with tube, is produced by the weight or pressure of the water without, and measure

extent of that pressure on the lower orifice of the tube.

The Rinstration extended to the case of Liquids lighter then Mercury.

80. Let there be four jars, each about four inches in diameter, and more than inches in height, severally occupied by mercury to the depth of shout two is In the axis of each jar, let a tube be placed, of about one inch and a half in d ter, and about one fourth taller than the jar, with both ends open, and the orifice under the surface of the mercury. On pouring water into the jars, the cury rises in the tubes, as the water rises in the jars; but the mercury rises as

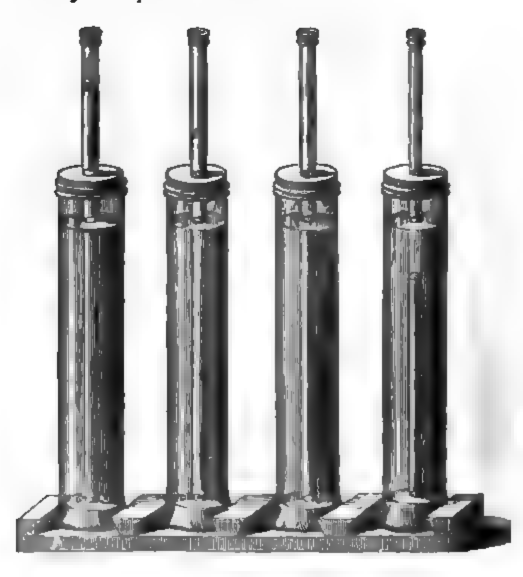
less than the water as it is heavier.

81. The mercurial columns in this case, as in the preceding experiment, owe existence to the pressure of the surrounding water, and by their height me the extent of that pressure on the areas of their bases respectively. They me considered as substituted severally for the aqueous columns, which would have tered the tubes had not the mercury been interposed. Accordingly, water poured into one of the tubes, the mercury in that tube subsides to a level wit mercury without, when the water poured into the tube reaches the level (water without.

62. The three remaining columns of mercury may be considered as substi in water, for columns of water, and being as much lower as heavier are found

quate to preserve the equilibrium.

83. It remains to be proved that other fluids, heavier or lighter than wester, w like manner be substituted for the columns of mercury, and of course for the we which the mercury is the representative.



17

14. Into the three tubes, in which, by the addition of water to the jars, columns of the part severally, ether, alcohol, (differently coloured, so that bey my be distinguished) and a solution of sulphate of copper, until the mercurial telem, within the tubes, are reduced to a level with the mercury without. It will be found that the column formed by the cupreous solution is much lower than the mire of the water on the outside of the tube; that the opposite is true of the tubes of alcohol; and that the other, still more than the alcohol, exceeds the surseming water in slevation.

CALORIC.

State water in elevation.

State in thus proved that columns of mercury, ether, alcohol, and of a size hand say, in water, be substituted for columns of this liquid; it is also appared that they must be as much higher as lighter, as much lower as heavier; or in

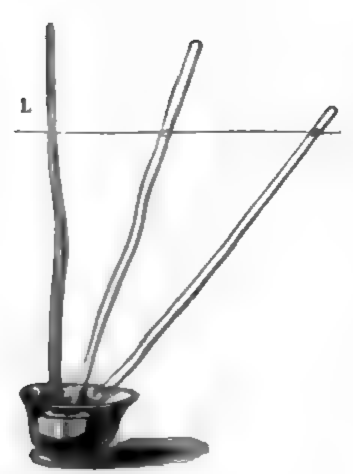
the week, their heights must be inversely as their gravities.

Torricellian Experiment.

The Pursuant is the law which has been above illustrated, that the pressure of one and may be adstituted for that of another, provided any difference of weight be repeated by a corresponding difference in height; if, in lieu of water, the mermy was presented by air on the outside of the tubes, unbalanced by air within, common of the metal would be elevated, which would be in proportion to the height

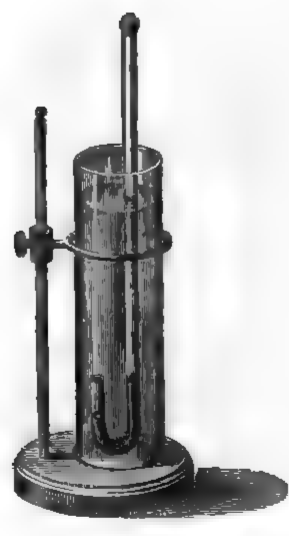
of vegit of the air thee acting upon it.

In order to show that the air exercises a pressure on the mercury outside of this, saulogous to that exercised by water in the experiments just described, it my request that this external pressure be unbalanced by the pressure of air this the tabe. This desideratum is obtained by filling, with mercury, a tube set three fact in length, open at one end and closed at the other, and covering the set with the hand, until it be inverted and merged in a vessel containing some the one metal, without allowing any air to enter. A mercurial column of about lacks in height will remain in the tube, supported by the pressure of the surming air, and an index of its weight. This is a case obviously analogous to it of the mercurial columns, supported by the pressure of water in the experiment destruction above given.



88. The tube may be supposed to occupy either of the three positions, represented in the drawing. The mercury, in each position, preserves the same degree of elevation, its surface being always in the same horizontal plane, or level, whether upright or inclined. Or see may suppose three tubes, filled with mercury, and inverted in a vessel, nearly full, of the same metal, to be placed in the positions represented in the drawing. The upper surfaces of the columns of mercury in each tube, will be found always coincident with the same horizontal plane, however different may be the angle which they make with the horizon. And the horizontal plane, in which their surfaces are thus found, will be between 25 and 31 inches above the surface of the mercury in the vessel The line, L, with which the increary in each of the tubes is on a level, represents a cord rendered horizontal, by mak. ing it parallel with the surface of the mercury in the reservoir.

dditional Rhotration of Atmospheric Pressure



80. I trust that the preceding illustra are well adapted to convey a clear co tion of atmospheric pressure; but as it: times happens, fortuitously, that when cannot get access to the mind ands form, it may reach it in another, even less eligible, I subjoin the following ill tion, which, though less amusing, and ciating with it fewer instructive phenos is more brief, and perhaps equally or sive.

90. If a tube, recurved into a crook : end so as to form a syphon, with is very unequal length, and both ends have the crook lowered into water, as adjoining figure, the fluid will of or without. But if, before the crook is in the fluid, it be occupied by mercur water will enter the tube, only so far pressure which it exerts upon the me in the short leg of the syphon, is compto raise the mercury in the long leg.

91. This pressure, or the effort of the to enter the tube, is obviously measure the height to which it forces the

the beight to which it forces the me in the long leg of the syphon, about mercurial surface in the short leg. height will of course be greater or h proportion to the depth to which the surface of the mercury may be sun will also be greater or less, according fluid in which it is immersed is hear

lighter. Hence, as water is about 820 times heavier than air, a depth of 630 . in air would displace the mercury as much as one inch in water.

98. Let us imagine a tube recurved at one end, similarly to the one represent the foregoing figure, the crook likewise occupied by mercury, to have the sriftee as completely above the atmosphere, as the critice of the tube is about water in the jar. The mercury, in the short leg of the syphon, thus situated, be evidently exposed to a pressure, caused by the air analogous to that sue from water, in the case of the tube, as already illustrated; and this pressure air would, as in the case of the water, be measured by the rise of the mercury

long leg of the syphon.

33. Yet to realize this experiment with a syphon reaching above the atmost section for giving such a height It is obviously impossible; but, as the only motive for giving such a height syphon is to render the mercury in the long leg inaccessible to atmospheric protein object can be etherwise attained, the phenomenon may be exhibited case of the atmosphere without any material deviation.

94. In fact, to protect the mercury in the long leg from atmospheric presses have only to seal the orifice of that leg, and, through the orifice of the other the syphon with mercury, before we place it in a vertical position. We shall find that the pressure of the air on the mercury, in the open leg of the sypho support a column of this metal in the other leg of nearly thirty inches, though sionally varying from 28 to 31 inches.

Inferences respecting the Weight of the Atmosphere from the preceding Emeric

25. Supposing the base of the column of mercury, sustained by the cinese as demonstrated in the preceding articles, were equivalent to a square inch, th weight of the column would be about fifteen pounds. This of course represen weight of that particular column of air only, whose place it has usurped; a for every other superficial inch on the earth's surface, a like column of air the earth must sustain a pressure from the atmosphere, equal to as many colu mercury, 30 inches high, as could stand upon it; or equal to a stratum of m of the height just mentioned, extending all over the surface of the globe.

19

It has been shown that the heights of heterogeneous fluids, reciprocally resisting another, are inversely as their gravities; or, in other words, that they are as much lower as heavier. The height of the column of an ward, by its pressure, elevates the mercury, must, therefore, be as much greater that he git of the column of mercury, as the weight of the mercury is greater than he weight of the air, supposing the air to be of uniform density. Mercury is liftimes neavier than air, and of course the height of the atmosphere would be in taken in density) 11152 × 30 inches = 27880 feet; supposing 30 inches to be height of the mercurial column supported.

Ecace the atmosphere, if of the same density throughout as on the surface of the extant would not extend much above the elevation ascribed to the highest moun-

But as the pressure of the atmosphere causes its density, it may be demonstrated that the heights increasing in arithmetical progression, the densities will burease in geometrical progression. Thus at an elevation of three miles, the air burg. It constrains, half as dense as upon the earth's surface:

A: 6	mile	es i	t w	ill b	e <u>1</u>	At 21	mil	es it	will	be	T 5	l a
y	•	-	•	•	Ĭ	_			-			_
12	-	-	•	•	i Ta				-			
					72				-			
					BI I						10	•

we meet that we can render it by the finest air pump. These results have been verified, to a considerable extent, by actual observation.

It is reasonable to suppose that there is a degree of rarefaction, at which the waget of the ponderable particles of the air will be in equilibrio with the repulsive power of the caloric united with them. Beyond the distance from the earth's surface at which there should be such an equilibrium, the air could not exist. Hence a saferred that the extent of our atmosphere is limited.

Of the Water Pump.

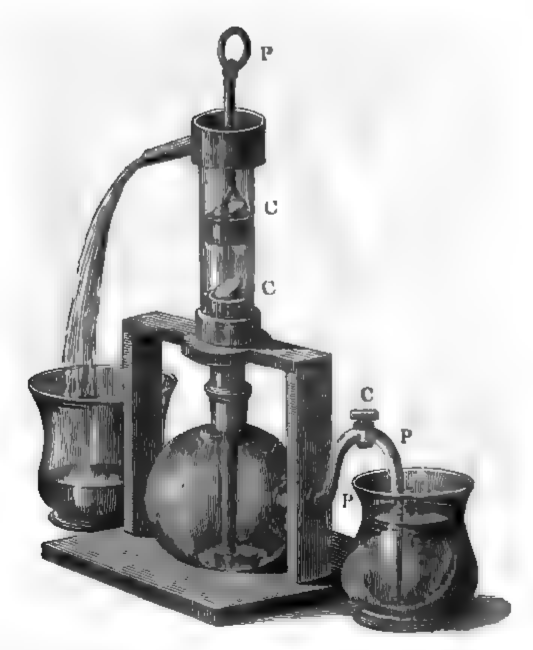
100. The admission of the atmosphere is necessary to the suction of the water than a receiver. Fir may be removed from close vessels by the same process. Water then the atmosphere; air presses out by its own elasticity.

Mechanism and Action of the Surtion Pump rendered evident by means of a Model with a Gam Chamber. Inference between pumping an Elastic Fluid and a Liquid, illustrate by an appropriate Contricunce.

A little suction pump is constructed, with a chamber C C, of glass, which person the action of its piston, P, and valves to be seen. Below the pump is a break green glade with water. This globe communicates with the pump by a took, ready descending from the lower part of the pump, through an aperture in the globe, the nearly reaches the bottom. This tube is luted air tight into the aperture ty where it enters the globe. Its orifice, next the chamber, is covered by a rare opening upwards. In the axis of the piston there is a perforation, also covered

by a vaire opening upwards.

ا المناهبة و المناهبة المناهب the as it race its valve will shut close; so that if nothing passes by the sides of the paling, por enters into the chamber of the pump from below, a vacuum must be Ispec behalf the piston. Under these circumstances, it might be expected that the water water in the from the globe through the lower valve, and prevent the forma-La :: a racum. But being devoid of elasticity, and, therefore, incapable of selfesterning beaund the space which it occupies, the water does not rise into the chamwer a the pump, so long as by means of the cock, C, of the recurved pipe, PP, comstartion with the external air is prevented. But if this cock be opened during Legisle movement of the piston, a portion of the water will mount from the game and the chamber at each stroke of the piston. The opening of the cock perma me at maphere to press upon the fluid in the globe, and to force it up the tube less as use pump chamber, as often as the chamber is relieved from atmospheric premara at the rise of the piston. As soon as the piston descends, the valve over the refer of the tube shuts, and prevents the water from returning into the globe. It is of course forced through the perforation in the piston, so as to get above it.



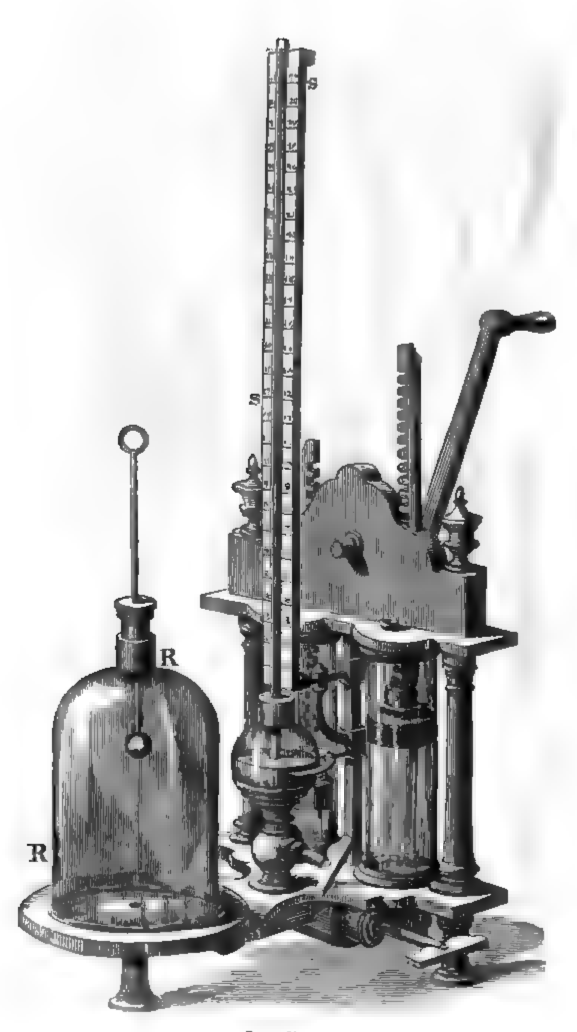
When the piston rises, the valve over its perforation being shut, it lifts the p of water above this valve until it runs out at the nozzle of the pump; while chamber, below the piston, receives another supply from the globe. But if after the water has been pumped from the globe, the pumping be continued with the closed, a portion of air will be removed from the globe at each stroke, until the

closed, a portion of air will be removed from the globe at each stroke, until the due be so much rarefied, as, by its elasticity, no longer to exert against the valueing the tube, sufficient pressure to lift it, and thus to expand isto the value formed behind the piston, as often as it rises.

103. The rarefaction thus effected in the air remaining in the globe, is rematrikingly evident, by causing the orifice of the curved tube to be under the set of some water in an adjoining vase, while the cock is opened. The water refrom the vase into the exhausted globe with great violence; and the extent e rarefaction is demonstrated by the smallness of the space within the globe while residual air occupies, after it is restored to its previous density by the entrar the water.

the water.

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(Page 21.)

CALORIC. 21

Description of a Chemical Implement.

104. The operation of sucking up a liquid through a quill, arises from the partial removal of atmospheric pressure from within the quill by the muscular power of the mouth. There is a great analogy between the mode in which suction is effected by the mouth, and that in which a liquid is made to rise into the bulb of an implement which I am about to describe, and which is very useful for withdrawing small portions of liquids from situations from which otherwise they cannot be removed without inconvenience.

105. This instrument is constructed by duly attaching a bag of caoutchook to the neck of a glass bulb with a long tapering

perforated stem.

106. In order to withdraw from any vessel into which the stem will enter, a portion of any contained liquid, it is only necessary to compress the bag so as to exclude more or less of the air from within it; then to place the orifice of the stem below the surface of the liquid, and allow the bag to resume its shape. Of course, the space within it becoming larger, the sir must be rarefied, and inadequate to resist the pressure of the atmosphere, until enough of the liquid shall have entered to restore the equilibrium of density between the air within the bag and the atmosphere. The air within the bag cannot, however, fully resume its previous density; since the column of the liquid counteracts, as far as it goes, the atmospheric pressure. Indeed, this counteracting influence is so great in the case of mercury, that the instrument cannot be used with

id. It is however the only substance, fluid at ordinary temperatures, which avy to be drawn up into the bulb of the instrument in question, when fur-

rith a stout bag.

Of the Air Pump.

Difference between the Air Pump and the Water Pump.

he action of the air pump is perfectly analogous to that of the water pump; as no difference between pumping water and pumping air, excepting that which om the nature of the fluids; the one being elastic, the other, in common with a general, almost destitute of elasticity.

n the air pump, as in the water pump, therefore, there is a chamber, and an id lower valve, which operate in the same manner as the valves of the water

ready described.

Description of a large Air Pump with Glass Chambers.

The opposite engraving represents a very fine instrument of large size, obfrom Mr. Pixii, of Paris.

From the figure, it must be evident that this pump has two glass chambers. we amountly large, being nearly three inches in diameter inside. The Ive, V. is placed at the end of a rod, which passes through the packing of Hence, during the descent of the piston, the friction of the packing the red. causes it to act upon the valve with a degree of pressure adequate at my escape of air, through the hole which it closes, at the bottom of the The air included between the piston and the bottom of the chamber, is, the descent of the piston, propelled through a channel in the axis of the evered by a valve opening upwards. When the motion of the piston is rethe air cannot, on account of the last mentioned valve, return again into the that the siston leaves behind it. But in the interim, the same friction of ring, about the rod, which had caused it to press downwards, has now, in same of the reversal of the stroke, an opposite effect, and the valve V is lifted a collar on the upper part of the rod will permit. The rise, thus permitted, fisient to allow the air to enter the chamber through an aperture which the desed, and which communicates by means of a perforation with a hole in med the air pump plate, and of course with the cavity of the receiver, RR, the plate. The reaction of the air in the perforation and pump chamber

being diminished, the air of the receiver moves into the chamber until the brium of density is restored between the two cavities. The chamber will now full of air as at first; but the air with which it is replenished is not so dense us I as the whole quantity in the receiver and the chamber scarcely exceeds that had existed, before the stroke, in the receiver alone. By the next downward a the air which has thus entered the chamber is propelled through the valve he the piston. Another upward stroke expels this air from the upper portion chamber; and the valve attached to the rod being again uplified, the portion chamber, left below the piston, is supplied with another complement of air flu receiver: and thus a like bulk of air is withdrawn at every stroke of the pussessy a like bulk of air, since the quantity necessarily varies with the density of in the vessel subjected to exhaustion. This density is always directly as the quarter remaining; of course it finally becomes insignificant. Thus when the quarter the receiver, is reduced to one-hundredth of what it was at first, the weight removed, at each stroke, will be one-hundredth of the quantity taken at each when the process began.

111. I have explained the action of one chamber only, as that of the other actly similar, excepting that while the piston of one descends, that of the

112. The gange represented in the engraving, is one which I have contrived a well known principle. It consists of a globular vessel to hold mersury, sup upon a cock. The mercury is prevented from entering the perforation in the by a tube of iron, surmounted by a smaller one of varnished copper, which pas take a Torricallian glass tube till it reaches near the top. The glass tube of its lower extremity, under the surface of the mercury in the globe. The exhaust take, and that of any vessel placed over the air nump plate, proceed si of this tube, and that of any vessel placed over the air pump plats, proceed si meously, and consequently the mercury is forced up from the globe into the tabe to an altitude commensurate with the rarefaction.

113. By inspecting a scale, 88, behind the glass tube, the height of the mis assertained. In order to make an accurate observation, the commencem the scale must be duly adjusted to the surface of the mercury in the globe. C account it is supported by sliding bands on an upright square bar, between the

cylinders.

114. The receiver, RR, represented on the air pump plate, is one which I t employ in exhibiting the artificial aurora borealis. The sliding wire, termina a ball, enables the operator to vary the distance through which the electrical cations are induced.

Experimental Illustrations of the Elastic Reaction of the Air.

Air occupying a small Portion of a Cavity, rarefied so as to fill the whole Spe 115. Air is dependent on its own weight for its density, and enlarges in b proportion as the space allotted to it is enlarged.



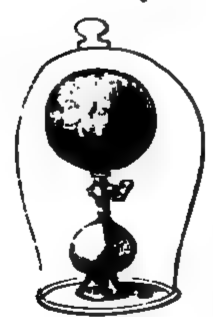
116. The mode in which the air occupying but i part of a vessel may be rarefied so as to fill the cavity, is shown by the experiment represented annexed engraving. A bladder is so suspended a vessel included in a receiver, as that the cavity bladder communicates through its own neck and the vessel, with the cavity of the receiver; whench communication exists between the receive the space between the bladder and the inside

117. Things being thus situated, and the receiv hausted, the bladder contracts in consequence removal of air from within it, proportionably wi exhaustion of the receiver; for, as the air between outside of the bladder, and the inside of the wat no longer resisted, within the bladder, by air of the density, it expands into the space which the b had occupied, so as to reduce it into a very a оопореля.

1(8. This cannot excite surprise, when it is recel that the air, confined between the outside of the

der and the inside of the vessel, had previously to the exhaustion been conapporting the whole atmospheric pressure, and must of course enlarge itself its elasticity, as that pressure is diminished.

Dutantion of a Coortchout Bug by the Rarefaction of confined Air.



119. The power of any included portion of air to extend itself in consequence of a removal of pressure, is illustrated in another way, by subjecting to a highly rarefied medium a gum elastic bag, its orifice being previously closed, so as to be air-tight. The bag will swell up in a most striking manner, in proportion to the diminution of power in the air without the bag to counteract the reaction of the air within it.

120. The experiment is reversed by subjecting a bag, while inflated, to the influence of a condenser, by which it may be reduced in size more than it had been expanded; the air within the receiver being rendered denser than without.

127. In the adjoining cut, the gum clastic bag is represented as when inflated. The glass represented below the bag, is one which happened to be used as a support when the drawing was made.



Expulsion of a Liquid by the Rarefaction of Air.

122. A flask, half full of water, is inverted in another vessel, having some water at the bottom, and both are placed, under a bell glass, on the plate of an air pump. As the bell is exhausted by the action of the pump, the air included in the flask enlarges its bulk, finally occupying the whole cavity, and partially escaping from the orifice through the water in the lower vessel. the atmosphere is allowed to re-enter the bell, the water rises into the flask, so as to occupy as much more space than at first, as the air occupies less, in consequence of a portion having escaped as abovementioned,



Experimental Proofs of the Weight of the Atmosphere. Atmospheric Pressure on the Hand.

123. If, as represented in this figure, the air be exhausted from a vessel covered by the hand, its re-moval will be found almost impracticable: for, supposing the opening which the hand closes to be equal to five equare inches, at 15 lb. per square inch, the pressure on it will evidently be seventy-five pounds.





124. Let there be a glass vessel open at both ends, as represented in this figure. Over the upper opening let a bladder be stretched and tied, so as to produce an air-tight juncture. For every square inch of its superficies, the bladder thus covering the opening in the vessel sustains a pressure of about 15 pounds. Yet vessel sustains a pressure of about 15 pounds this is productive of no perceptible effect; because the almosphere presses upwards against the lower surface of the bladder, as much as downwards upon the upper surface. But if the vessel be placed upon the plate of an air pump, so that, by exhaustion, the atmosphesic pressure downwards be no longer counteracted by its pressure upwards, the blad- x

der will be excessively strained, and usually tern into pieces.

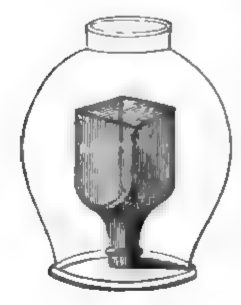
125. When the bladder is too strong to be broken by the unassisted weight of the sir, a slight score with the point of a penkuife will cause it to be ruptured not enly where the score is made, but in various other parts, so that it will, at times, be torn entirely from the rim of the vessel.

The Homispheres of Otho Guericks, the celebrated Inventor of the Air Pump.



126. Two brass hemisphers are so ground to fit each oth at their rims as to form an airtight sphere when united. O of the hemispheres is furnish with a cock, on which in screw for attaching the whole to the air pump. Being by

these means exhausted, the cock closed, and the ring, R, screwed on to the cock great force must be exerted, before the hemispheres can be separated.

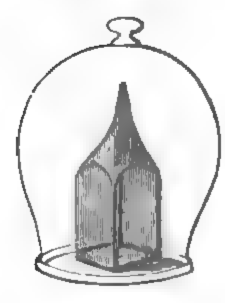


Battle broken by Exhaustion of the Air from within.

127. Proof that a square glass bottle may be broken by atmospheric pressure on the outside, as soon as it courses to be counteracted by the resistance of the air within.

128. The mouth of a square bottle being placed over the hole in an air pump plate, so as to be sufficiently tight for exhaustion, a few strokes of the air pump, by withdrawing the air from the interior, causes the bottle to be crushed.

129. A stout globular glass vessel, with an aper-ture at top, is placed over the bottle, to secure the spectators from the fragments.



Bottle broken by Exhaustion of the Air from without.

130. The elastic reaction of the air, confined within a square bottle, will burst it, as soon as relieved from the counteracting weight of the atmosphere.

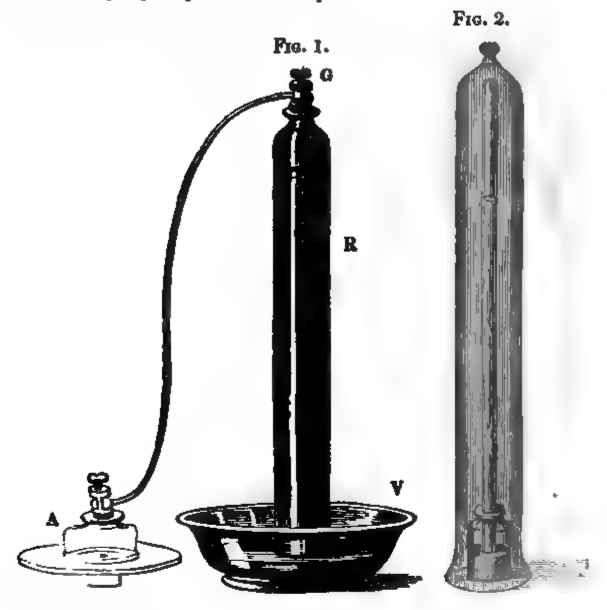
131. If a thin equare bottle, so sealed that while unbroken the contained air cannot escape, be placed within the receiver of an air pump, the exhaustion of the receiver will, by removing the pressure which counteracts the clastic reaction of the confined air, cause the bottle to be fractured.

The Height of the Column of Mercury which belunces the Atmosphere, shown by Ezhaustion.

132. R, fig. 1, is a bollow glass cylinder, about 33 inches in height, and 24 inches in diameter, into the upper end of which a brass gallows screw, G, is comented; so that by means of the flexible pipe communicating with the air pump plate, A, the cylinder may be exhausted. The mouth of the cylinder being immersed in mercury in the vase, the metal, as the exhaustion proceeds, ruses in the cylinder, until #

25

conver less mearly to the height at which it stands in a Torricellian tube, into pump may be more or less perfect.



Berometric Column of Mercury lowered by Exhaustion.

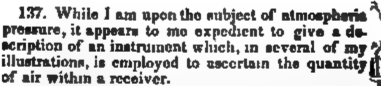
123. It has been shown that in a tube word of air, a mercurial column may be supred a the height nearly of thirty inches; and this has been alleged to result to pressure of the atmosphere on the surface of the mercury on the outside of a take.

134. In order to verify this allegation, let a tube, fig 2, supporting within it a bean of mercury, he placed under a competent receiver upon the air pump plate.

13. It was be found that, as the air is withdrawn from the receiver, the mercury for tube will subside, and, if the exhaustion be carried far enough, will sink to a start the mercury on the outside.

If, while this experiment is performing, a communication exist between the pump and the cylinder, R, employed in the preceding experiment, the mercury last is cylinder, while it falls in the tube, thus proving that the force which there is the cylinder are from the outside of the tube and lower the increase hast is adequate to raise in the cylinder a mercurial column equal in height to twich a reduced.

Of the Barometer Gauge.



13d. It consists of a barometer tube, 33 inches in height, supported in a vertical position by a peda tal, and a strip of wood, G.G. Attached to the latter is a brass scale, by which 30 inches is divided into 500 equal parts. The gauge tube is surmounted by a ferrule and gallows screw, by the aid of which a flexible leaden pipe, P, communicates with the bore of the tube. By means of the valve cock and gallows screw at V, this pipe may be made to communicate also with the cavity to be measured, the valve took enabling us to suspend the communica-tion when desirable. The lower orifice of the glass tube, T, is covered by morcury in a broad shallow receptacle, D. Supposing the cavity, under these circumstances, to be exhausted, and the communication with the bore of the glass tube open, the extent of the exhaustion, or, in other words, the quantity of air withdrawn, will be exactly in portion to the rise of the mercury as indicated by the scale; and consequently, reversing the operation, the fall of the mercury, as indicated by the scale, will show the quantity of air which may be introduced. If we count the degrees upwards from the surface of the mercury in the receptacle, D, their number will show the quantity of air withdrawn. If we count the degrees downwards from the level of the top of the mercurial column in the barometer, the number will indicate the exact quantity of gas in the cavity examined. In short, the quantity taken out, or introduced, is always measured by the number of degrees which the mercury rises or falls in

consequence. It is preferable to have two scales, one beginning above, the other below.

G

130. This gauge may be employed to indicate the quantity of air in any cavity. It only requires accuracy in the divisions of the scale, and in the adjustment of zero to the proper level. As the height of the mercurial column in the barometer varies with those changes of atmospheric pressure which it is employed to indicate, therefore, in counting downwards, care must be taken to place the commencement of the scale on a level with the upper end of a column of increury in a good barometer, at the time. To facilitate this adjustment, I have occasionally placed a Torricellian tube by the side of the gauge tube. The top of the column of the increury in the Torricellian tube is then the proper point for the upper zero. As the strip of wood to which the scale is attached slides upon the iron rod, R, the scale may be fixed at a proper height by a set screw."

140. As a perfect vacuum cannot be produced by means of an air pump, in order to wash out of a receiver all traces of atmospheric air, it is necessary that portions of the gas to be substituted should be repeatedly introduced, and as often removed by exhaustion.

141. The rise of the mercury in the tube, by diminishing the quantity in the receptacle, D, will cause the surface of it to be lower; but the breadth of this vessel is so great, and the descent of the mercurial surface in it so inconsiderable, that no error worthy of attention is thus produced.

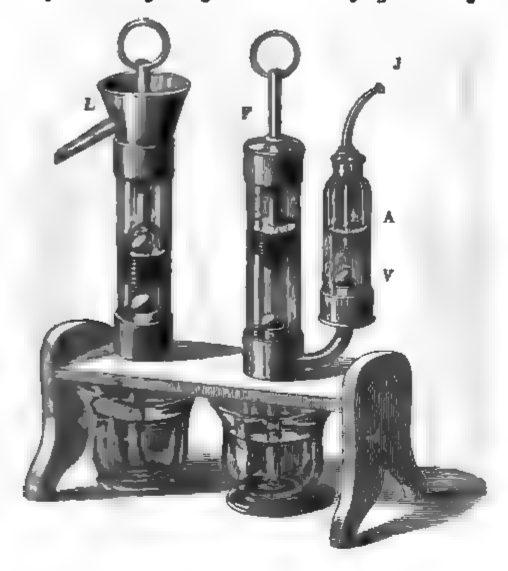
142. It is proper to mention that the cavity of the tube ought to be so small in proportion to that of the receiver, as to create no error worthy of attention.

^{*} Both the gauge tube and the rod, R, should be longer than they are represented in the figure.

t One gas may be employed to wash another out of a cavity, in a mode analogous to that in which water may wash out alcohol, or alcohol water

caloric. 27

Aparetus for illustrating the Difference between the Lifting and Forcing Pumps.



IC The process by which the water is drawn into the chamber is the same in some of the forcing as in that of the lifting pump. In the lifting pump, L, the ster which has entered the chamber during the ascent of the piston, passes through a panel of iring its descent, and is lifted by it when the motion is reversed. In the range panels F, the piston, being imperforate, forces, in descending, the water into marketing air vessel. A, whence its regress is prevented by a valve, V. The water repealed, the water accumulates in the air vessel, compressing the contest ar analyst reacts upon the water with sufficient force to cause an emission of water through the jet pipe, J J, commensurate with the supply.

Of Condensation.

16. It has seen shown that, in consequence of the elasticity of the air, the quanty of the field, in any close vessel, may be diminished until the residual portion a by the artion of the air pump, become too rare to escape in opposition to the ty algebrase entack made by the valves. It remains to show that, in consequence has same property, by an operation the converse of that of the air pump, the air my adequate vessel may be made many times more dense than it would otherwise.

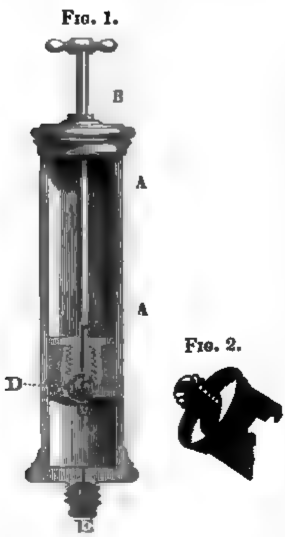
Of the Condenser.

15 The instrument employed for the purpose of condensing air is called a con-

The air pump was illustrated by its analogy with the suction pump. There was analogy between the condenser and the forcing pump. In the air pump, which between the chamber and receiver opens towards the chamber; in the case the makener a corresponding valve opens towards the receiver.

budes the valve thus placed between the chamber and receiver, there is in the partner valve. In the air pump, the air passes this second valve only

when the piston moves so as to been the vacancy between it and the botchamber, in the condenser, the air passes only when the piston moves a large the vacancy. In other respects these machines are so much alike, the might be used for the other. In my experimental illustrations, I shall have to employ instruments which serve either to exhaust or to condense, at the aperture selected for making a communication with the receiver.



represents a condenser. It cobrass cylinder, A A, ground intense to be perfectly cylindrical. a piston, B, is fitted by mean leathers packed between scresented in the figure, and turn lathe, so as to enter the chambe ence to considerable force. At end of the rod, a perforation, C seen, which commences at the tremity, rises vertically until it the packing, and then passes o angles to its previous direction t rod of the piston. Just above commences, a cavity, D, may be which is left for the upper vivalve is formed of a strip of oi tied over a brass knob represent the cavity.

149. The upper and lower valuetly alike; hence, a good ides may be obtained from fig. 2, what a separate view of the lower value 150. The action of the conditions. When the piston is dritted air within the chamber gets packing through the perforation the upper valve, which opens

with ease so us to afford a passing the piston descends, the air inch chamber cannot get by the leath

The upper valve at the same time shuts so as to prevent it from getting t perforation, C.C. It has therefore to proceed through the lower perf. The piston being drawn up again, the valve at E shuts and prevents a reair expelled, while the air of the chamber again gets below the piston as instance. Thus, at every stroke, the contents of the chamber are discharg the lower valve, while its retrocession from any receiver into which it is prevented by the valve, E.

151. As the quantity of air in the vessel increases, the force requisite t piston home becomes greater; and it has to descend farther, ere the air chamber exceeds in density that in the receiver, so far us to open the low

Influence of Pressure on the Bulk of Air, and of its Density on its Ren

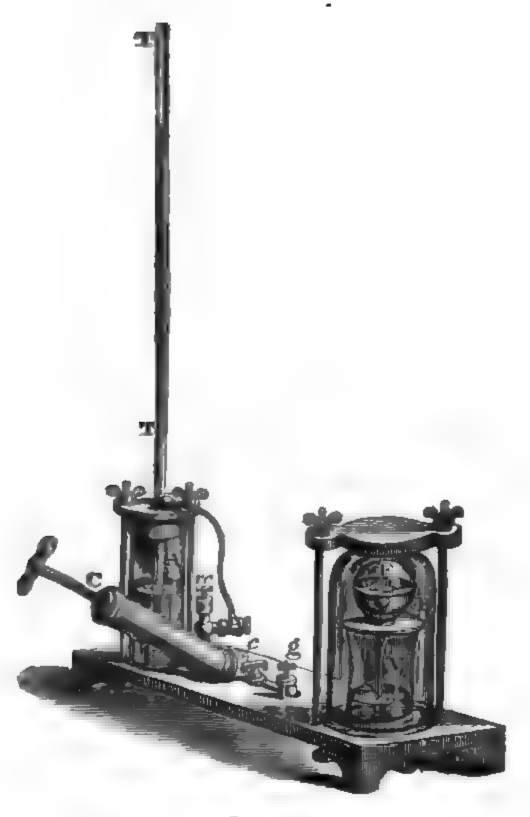
152. Air lessens in bulk as the pressure which it sustains augments; and ance arising from its elasticity is augmented, as the quantity confined in the is increased, or the confining space diminished.

153. For the illustration of this proposition, I have devised the appar

sented in the opposite engraving

154. If mercury be poured into the air-tight vessel, A, through the which passes perpendicularly into this vessel until it touches the bottom in the vessel cannot escape, it is gradually reduced in bulk, but at the sar acts upon the surface of the metallic liquid with a force which becomes proportion as its bulk is lessened. Hence an increasing mercurial columpheld, which by its height indicates the resistance. When the air in has been reduced to half its previous bulk, the height of the mercury in the about 30 inches, or equal to that of the mercury in the barometer at

rates for illustrating the Influence of Pressure on the Bulk of Δir .



(Page 28.)

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29 CALORIC

Thus it is shown, that when air is condensed into half which it occupions under the pressure of the atmosphere, its reactive power, being adequate to support a column of mercury equal to the pressure of them, in addition to that pressure. It follows that the quantity of air octay space is as the pressure, and is always to that of an equal bulk of the a, as the height of the column of mercury which the said air can support ellies take, in to the height of the mercury in the barometer; and like-the resistance of air increases with the diminution of the including space; and that the space which a given weight of air is excellent of conveying s, that the space which a given weight of air is espable of occupying, Pressure increases

us to be shown that the resistance of air to compression increases as

ty is any space increases.

by means of the condensar, C, (the valve cook, e c, and the cook, c, being the injusted into the vessels, A and B at the same time, it will be found liquid in the vace, V, will mount into the flack, F, and that when the president to cause the air in this to be reduced to half its previous volume, my is the tube, TT, will have the same height as in the previous experiment the density of the sir, and of course its quantity and reactive power, at in one case no less than in the other.

r currentication but ween the condenser and the receiver, A, is suspended a feat mentioned experiment, by closing the valve cook, v.c. This cook duing the action of the condenser in the second experiment; and likeg the action of the condenser in the second experiment; and like-sch at c, which corves to intercept the communication between the

ar and the receiver, B.

Sichemical Action of the Lange in Respiration illustrated.

The chretica of the storage rareles the air within the cavity of the thorax. mostly, the atmospheric pressure not being adequately resisted, the external factoring the traches into the lungs, dileting all the cells. The depression summand consequent diminution of the cavity cause the air which had wind, at an equivalent portion, to flow out. For the illustration of the probable contributed, I have contrived the apparatus represented below.

3. Itali receiver, R., with an orifice, O, is placed in a globe containing water,





so that about two-thirds of the receiver are occupied by this liquid, the rem with air, whilst a bladder is so suspended from the orifice as not to touch the

160. The atmosphere has access to the cavity of the bladder through its nech through the orifice O of the receiver, but not to the space A, between the outs the bladder and the inside of the receiver.

161. It may be assumed as an obvious consequence of the preceding experience (154, 156) that the pressure, exerted by any given quantity of air, is inversely confining space; or in other words, that the pressure increases as the space le

and diminishes as the space enlarges.

162. When a cavity to which the atmosphere has no access is enlarge density of the contained air is proportionably diminished. When any cavity minished, the density of the contained air is proportionably increased. But atmosphere, meanwhile, have access to the cavity, it will by its influx or efflut to preserve the equilibrium of density and pressure between the air of the cavithe external medium. These consequences are well known to ensue, from ternate enlargement and diminution of capacity, during the working of an air a condenser, or bellows.

163. In like manner the elevation of the receiver, R, enlarging the cavity it unoccupied by water, causes the air to rush in through the orifice, O; and t versal of the motion, reducing the cavity, causes the air to rush out through same aperture. The bladder is so situated as to receive all the air that enters, supply all that is expelled. Hence when the receiver is lifted, the bladder flated, and when lowered to its previous position, the bladder resumes its or

dimensions.

164. Supposing the space, A, between the outside of the bladder and the intereceiver, to represent the space between the outside of the lungs and the of the thorax, the cavity of the bladder representing the cavities of the lung the orifice, O, performing the part of the trachea and nostrils, the explanation, given, will be as applicable to the apparatus by which nature enables us to be as to that employed in the preceding illustration.

EXPANSION OF ELASTIC FLUIDS.

165. Having by means of the preceding digression plained the nature and extent of atmospheric pressushall proceed to show the important influence exercity it in all chemical processes in which elastic fluids concerned.

166. It has been demonstrated (54) in illustrating principle of Sanctorio's thermometer, that the bulk of air in any space varies with the temperature.

167. It has been shown that the same effect ma produced by variations in atmospheric pressure. (

119, 120, 122.)

- 168. It follows that the volume of elastic fluids is versely as the pressure and directly as the heat. In c words, the less the pressure and the greater the heat, larger their bulk; and vice versa, the less the heat, the greater the pressure, the less their bulk.
- 169. Agreeably to the observations of Dalton, Gay-Lussac, and (ton, 1000 parts of atmospheric air, in rising from the temperature of to 212°, will expand so as to measure 1375 parts nearly, or, $\frac{1}{480}$ th obulk which it would have at 32°, for each degree of heat which it makes it.

CALORIC. 31

Having, therefore, any given bulk of dry air, 100 cubic inches for 121 60°, to find its bulk at any other temperature, suppose at 80°, in the first place consider that 480 parts at 32° would at 60°, no part for every degree above 32°, be 508 parts; and would by tionate increase, become at 80°, 528 parts. But if 508 parts at me 528 at 80°, what will 100 parts at 60° become when heated

508:528::100:103.9

It has been inferred by the same distinguished philosophers, that prin substances, whether gases or vapours, are expanded by heat at a rate as dry atmospheric air, if they be not in contact with any able matter, in the liquid or solid state, which by vaporizing or contact wary the result.

Theory of Expansion.

The expansion of matter, whether solid, liquid, or aëriform, by an enterperature, may be thus explained.

In proportion as the temperature within any space is raised, there more caloric in the vicinity of the particles of any mass contained pace. The more caloric in the vicinity of the particles, the more il combine with them; and in proportion to the quantity of caloric mbined, will they be actuated by that reciprocally repellent power, in proportion to its intensity, regulates their distance from each

There may be some analogy between the mode in which each able atom is surrounded by the caloric which it attracts, and that in the earth is surrounded by the atmosphere; and as in the latter case, as in the inner, the density is inversely as the square of the dis-

At a height at which the atmospheric pressure does not exceed a the square lach, suppose it to be doubled, and supported at that inspective by a supply of air from some remote region; is it not a that a condensation would ensue in all the interior strata of the pressure would be doubled throughout, so as to be the terrestral surface, 30 pounds, instead of the pressure pressure in pressure at the point from which the change would be and I would not exceed two grains per square inch.

6. In the timer, it may be presumed that the atmospheres of caloric streets in quantity and density about their respective atoms, by a

incress in the caloritic tension of the external medium.

'honical Action, where Elastic Fluids are to be generated recolved.

Of Vaporization.

Water would boil at a lower temperature than the atmospheric pressure was lessened; for when it

has ceased to boil in the open air, it will begin to boil again



in an exhausted receiver. Those who ascend mountains find that for every 530 feet of elevation, the boiling point is lowered one degree of Fahrenheit's thermometer. It is, in fact, lowered or raised what has a degree for every tentl of an inch of variation in the height of the mercury in the barometer.

Ebullition from diminished Pressure.

178. The adjoining figure represents a vessel o water boiling within a receiver, in consequence o the diminution of pressure by exhaustion.

Culinary Paradox.—Ebullition by Cold.



179. A matrass, half full of water, be ing heated until all the contained air i superseded by steam, the orifice is closs so as to be perfectly air-tight. The mateur is then supported upon its neck, in an in verted position, by means of a circula block of wood. A partial condemnation of the steam soon follows from the re frigeration of that portion of the which is not in contact with the water The pressure of the steam upon the liqui of course becomes less, and its boiling point is necessarily lowered. Hence i begins again to present all the phenomes of challition, and will continue boiling sometimes for nearly an hour-

1×0. By the application of ice, or of sponge soaked in cold water, the ebullition is accelerated; because the aqueous vapous which opposes it, is in that case more re-

pidly condensed; but as the calorie is at the same time more rapidly at structed from the water by the increased evolution of vapour to replace that which is condensed, the boding will cease the sooner.

Ingreed Apparates for showing the Odinary Parades.

181. This figure illustrates a new and instructive method of effecting ebullition by cold.

162. The apparatus consists principally of a glass matress, with a neck of about three feet in length, tapering to an orifice of about a quarter of an inch in diameter. The bulb is bulged inwards in the part directly opposite the neck, so as to create a cavity capais of holding any matter which it may be desirable to have situated moreons holding a fion to the matress, a receptacle holding a few pounds of mercury is requisite. bulb of the matress being rather less than balf full of water, and this being heated to ebullition, the orifice should be ejesed by the finger, defended by a piece of gam-blas-tic, and depressed below the surince of the mercury; the whole being supported as re-presented in the figure. Under these circumstances, the mercury rises as the temperature of the water declines, indicating the consequent diminution of pressure within the tulb. Meanwhile, the decline of pressure lowering the boiling point of the water, the ebullition continues till the mercury rises in

stak meanly to see height of the mercury in the barometer.

68. By introducing into the cup formed by the bulging of the bulb, cold ex, alcohol, ethar, or ice, the refrigeration, the diminution of pressure, is a chulision, are all simultaneously accelerated; since these results suprocally dependent on each other.

primental Proof that some Liquids would be permanently aëriform, if Atmospheric Pressure were removed.

184. The power of certain liquids, common ether for instance, to assume in vacuo, at ordinary temperatures, the aëriform state, in opposition even to the pressure of a column of mercury, may be shown by

185. A glass funnel is ground to fit air-tight into the neck of a glass decanter, so that the stem of the funnel may reach nearly to the bottom of the decanter, as represented in the adjoining cut. The decanter is filled with mercury, with the exception of a small portion of the neck, which is occupied by other. The stem of the funnel is then introduced into the neck of the decanter, so as to be air-tight; and the whole being included in a receiver, the air is withdrawn by a pump. The other converted into vapour will force the mercury to rise from the decanter, through the stem, into the wider part of the funnel.

Retionale.—The attraction between the ponderable particles of the latter; since one tends to rarefy the caloric,

the other to condense it into the limited space occupied by the ethe follows that the caloric cannot combine with the ponderable matter to the point at which the repulsive power becomes equal to the attractive the repulsion exercised by the same number of particles of caloric greater as the space is less, and vice versa. The larger, therefore space occupied by the ponderable particles of the ether, the more may combine with them, without rendering its reciprocally repulsive paramount to its attraction for them.

187. The removal of atmospheric pressure, by allowing the pone particles to occupy a larger space, enables them to combine with that tional quantity of caloric which is necessary to the aëriform state.

188. This explanation may, of course, be extended to the ebullit other liquids in vacuo, at temperatures lower than those at which boil in the air. It is obviously applicable to the two preceding il tions.

Boiling Point elevated by Pressure

189. Into a small glass matrass, with of about an inch and a half in diameter neck of about a quarter of an inch in be troduce nearly half as much ether as wo it. Closing the orifice with the thumb, ho bulb over the flame of a spirit lamp, ur effort of the generated vapour to escape be difficult to resist. Removing the matra sufficient distance from the lamp, lift the from the orifice. The ether, previousl escent, will rise up in a foam, produced rapid extrication of its vapour.

190. This experiment may be performed less risk, by plunging the matrass in hot instead of heating it by a lamp.

191. Having supplied a small flask with a quof mercury, sufficient to cover the bottom to at inch in depth, let there be a glass tube so intruthrough the neck, and luted air-tight, as to nearly an inch below the mercurial surface. flask thus prepared, be duly heated, the ether proportionably vapourized, and the generated pressing on the mercury, will cause a column metallic liquid to rise within the tube, and thus dicate and measure the pressure. It is necess discontinue the heat, when the mercurial coluproaches the upper orifice of the tube, in order vent the metal from overflowing.



High Pressure Boiler.



(Page 35.)

High Pressure Boiler.

M. That the temperature of steam increases with the pressure, may be described by means of a small boiler, such as is represented in the temperature.

IM. A gines tube, of about five feet in height, and of half an inch is her mendy, is accured into an aparture in a very strong iron boiler, an a to be air-tight, and so as to be concentric with the axis of the boiler. Within the boiler the tube descends in such manner as to pass through the star with which it is supplied, and to terminate close to the bottom, he much a small quantity of mercury purposely introduced. On the apposite with of the boiler, a tube, not visible in the angraying, descends into it. This tube commits of about two inches of a manifest harred, and is closed at bottom. The abject of it is to contain some mercury, into which the bulb of a thermometer may be plunged for acceptaining the temperature.

124. When the the been applied during a entire out time, the marcular will are in the glace tube so so to be visible above the beiler; and committee in the during the application of the five, it will be found that, with twent searche increment in its height, there will be a corresponding rise of

be succery in the thermometer.

186. In front of the tube, as represented in the figure, there may be obtained analy valve with a lever and weight for regulating the presented has been found that, when the effort made by the steam to escape, in the interest thus loaded, is equal to about fifteen pounds for every space inch in the area of the aperture, the height of the column of mercay, CC, raised by the same pressure, is about equal to that of the column of this metal, usually supported by atmospheric pressure in the tube of a brometer.

Hence the boiler, under these circumstances, is conceived to sustain whilanced pressure equivalent to one atmosphere; and for every additional filters pounds per square inch, required upon the safety valve to restain the steam, the pressure of an atmosphere is alleged to be added. To give a steam at 212 degrees, or the boiling point, such an augmentation of pour, a rise of 38 degrees is sufficient, making the temperature equal to 250 degrees. To produce a pressure of four atmospheres about 293 degrees would be necessary. Right atmospheres would require nearly 343 degrees.

197. When by means of the cock an escape of steam is allowed, a cor-

reporting decline of the temperature and pressure ensues.

196. If the steam, as it issues from the pipe, he received under a portion of water of known temperature and weight, the consequent accession of the in surprisingly great, when contrasted with the accession of weight dived from the same source. It has in fact been ascertained that one makes of water, converted into aqueous vapour, will, by its condensation, mise about ten measures of water in the liquid form one hundred these.

If the Incompetency of a Jet of High Steam to scald at a certain Distance from the Aperture.

160. Much attention has been excited by the observation, that the hands because boiler without

inconvenience, at a certain distance from the aperture through wheescapes.

Experimental Demonstration.

200. The fact that the hand may be immersed wit injury in a jet of steam while issuing from a boile not too near the aperture, experimentally demonstrate

201. Rationale.—Since the temperature, density, and pressure, form the distinguishing attributes of high steam, cannot be sustained out confinement, steam ceases to be high steam as soon as it is libe Consequently, a jet from a high pressure boiler is essentially no more a copious jet of aqueous vapour at the heat of boiling water.

202. The only distinguishing characteristic, derived from its prevenue and density, is a greater velocity of efflux. We any superiority of temperature, the high pressure jet is propelled in atmosphere with a momentum, which cannot be given to low steam. It the rapid refrigeration to which the former is subjected, at a sufficient tance from the place of its efflux to admit of an extensive diffusion atmosphere.

Ribustration of the Process by which Thermometers are supplied with the Liquic in their construction.



203. A globe, with a long cylindrical neck, situated as in the preceding and containing a small quantity of water, being subjected to the flame of a the water, by boiling, will soon fill the cavity of the globe and neck with When this is accomplished, bubbles of air will cease to escape from the orifice neck through the water in the vase.

204. The apparatus being thus prepared, on removing the lump, the wat quickly rush from the vase into the vacuity arising from the condensation

steam within the globe.

Explosive Power of Steam.

205. If a glass bulb, hermetically scaled while containing a small quantity of water be suspended by a wire over a lamp flame, an explosion soon follows, with a violence and noise which are surprising, when contrasted with the quantity of water by which they are occasioned.

206. Rationale.—Supposing that the bulb were, in the first instance, merely filled with steam, without any water in the liquid form, the explanation of this phenomenon would be comprised in the theory of expansion, already suggested. (178.) In that case, the effort of the steam to enlarge itself, would be nearly in direct arithmetical proportion to the temperature; but water being present in the liquid form, while the expansive power of the steam, previously in existence, is increased, more steam

rated with a like increased power of expansion. It follows that the mats of heat being in anthmetical proportion, the explosive power of fined vapour will increase geometrically, being actually doubled as

mas the temperature is augmented 38°. (196.)

rating Inperiments with respect to Vaporisation under extreme Pressure, by M. Cagniard de la Tour, and Mr. Perkins.

187. Agrembly to some experiments performed by M. Cagniard de la ur, is which liquids were exposed to heat in very stout tubes, vaporizawe performed in a space which was to that previously occupied,—

Ether, as 2 to 1, producing a pressure of 83 atmos-

Alcohol, as 8 to 1, producing a pressure of 119 atmospheres.

Water, as 4 to 1, producing a pressure greater than that caused by the alcohol.

B. Mr. Perkins alleges that a small iron boiler of great strength may be heated bet while holding a portion of water, and that if, under these circumstances, an ture be spened of 1 of an inch in diameter, the steam will not escape, although a soduction of temperature, it will rush out with great violence.

L It was inferred that the repulsion between the particles of the caloric in a with the water, and those in union with the metallic ring bounding the aperwas paramount to the pressure tending to produce the expulsion of the steam. Liam mable to reconcile this experiment with one which I performed by ngto insundecence, in a forge fire, a tube of iron, of which the bore was less for an inch, while, by means of a cock, a communication with a high pressure rwas made. Under these circumstances, the steam was not prevented from through the pipe.

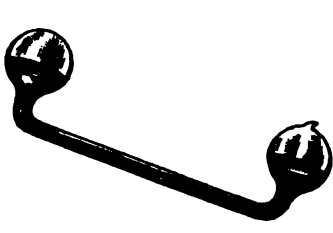
L it appears to be sufficiently proved that the quantity of caloric combined with weight of steam is always the same, whatever may be its temperature; the We have increasing and the latent heat diminishing as the density and pressure

Nymental.

Cold and Cloudiness arising from Rarefaction.

Import rarefaction in the air of a receiver is usually indicated by a cloud, when the exhaustion has proceeded beyond a certain point. A memoter placed in the receiver, shows that a decline of temperature accompanies this phenomenon. We may, therefore, infer that the cloud is the consequence of refrigeration. If the suggestions be correct which were made (Theory of Expansion, 175) respecting the mode in which caloric exists in atmosphere around the particles of ponderable matter, it will not be difficult to understand why acriform fluids should absorb more caloric, in proportion as their constituent particles are enabled, by a diminution of pressure, to become more remote Hence, by rarefaction, the capacity of air is increased, and cold is produced, which condenses the aqueous vapour until its sensible heat is restored by an accession of caloric from the surrounding medium. (184.)

Cold produced by the Pelm Glass.



213. In forming the bulbs severally at the ends of the glass tube represented in this figure, one is furnished with a perforated projecting beak. By warming the bulbs, and plunging the orifice of the beak into alcohol, a portion of this liquid enters, as the air within contracts in returning to its previous temperature. The liquid, thus introduced, is to be boiled in the bulb which has no beak, until the whole cavity of the tube and of both bulbs, not occupied by liquid alcohol, is filled with its steam. While in this situation, the end of the beak is to be shortened and scaled, by subjecting it to the flame excited by a blowpipe.

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214. As soon as the instrument becomes cold, the steam, which had filled the space vacant of liquid alcohol, condenses, and with the exception of a slight portion of vapour, which is always emitted by liquids when relieved from atmospheric pressure, a vacuum exists within the bulb.

215. The instrument, thus formed, has been called a palm glass; because the phanomenon which it exhibits is seen by grasping one of the bulbs, so as to bring it completely into contact with the palm of the hand. One of the bulbs being thus situated, and while surcharged with the alcohol, and held in the position represented in the figure, both the liquid and vapour are propelled from it into the other bulk. This phenomenon combines the characteristics of the differential thermometer, (69) with those of the culinary paradox, (179,) being the joint effect of the expansion, and evolution of vapour, in one part of the apparatus, and its contraction and condensation in another. The phenomena are precisely similar, whether we warm the lower bulb, or cool the upper one by means of ice. The motive for recurring to the experiment here is to state that, as soon as the last remnant of the liquid is forced from the bulb in the hand, a striking sensation of cold is experienced by the operator.

216. This cold has been attributed generally to an increase of the capacity of the residual vapour for caloric in consequence of its attenuation. The analogy is evident between this phenomenon and that above described, as taking place in the receiver of an air pump; in either case refrigeration results from a diminution of density.

Cold consequent to relaxation of Pressure.

217. Cold is produced whether a diminution of density arise from relieving condensed air from compression, or from subjecting air of the ordinary density to rarefaction. A cloud similar to that which has been described as arising in a receiver partially exhausted, may usually be observed in the neck of a bottle recently uncorked, in which a quantity of gas has been evolved in a state of condensation by a fermenting liquor.



i with a test k bas en air t , after due e will app , in the colar rays, to proeclosm. At the statementer will show that the on is productive of warmily,

arise (ER.) in air suddenly seen cares inches a y, no Mustrating's motocralegical , which he considers as the prin se of min sterms. This induced s, which he con to make some expert displate this subject.

the containing about a subject foot of space, furnished with the term, were made to communicate, respectively, with revenir, and of air regists with aqueous vapour." The sold, way decree of confection to account the selection of the sold. my degree of rarefaction, appeared to be the same, whether state or the other; provided that the air, replets with aqueous stact with Equid water in the vessel subjected to exhaustion. exptien of calorie, the cold produced was nearly twice as great as a not in contact with liquid water; being nearly as 9 to 5, as circumstances last mentioned, the hygrometer was motionless; to liquid water was accessible, the space, although previously extense, by the removal of a portion of it together with the air which is a sylvantion, accretize a capacity for more wanted. e exhibition, acquires a capacity for more vapour; and hence the metatraction of eas-third of the six, revolved more than sixty de-But when a smaller receiver (after being subjected to a me of about ten inches of mercury, so as to cause the index of r to more about thirty-five degrees towards dryness) was surrounded nixture, until a thermometer in the axis of the receiver stood at three finaling, the hygrometer in the axis of the receiver stood at three finaling, the hygrometer revolved towards dampness, until it went me layend the point at which it rested when the process commenced, in, therefore, that the dryness produced by the degree of rarefaction we than counterbalanced by a freezing temperature, acts the heat imported to the air above mentioned, the flet, that the gation in the case of air replate with vapour, and in that of ankyously great, and that when water was present the cold was greater and, led to the idea, that the heat arising under such circumstances would be different in angular the heat arising under such circumstances. ench efficacy in augmenting the buoyancy of an ascending column them, by an appropriate mechanism, the refrigeration was measured by af pressure at the moment when the exhaustion was arrested, and moments had become stationary, it was found exteris paritus, that the pressure arising from cold was at least one-half greater in the anhymis in the air replete with vapour. This difference seems to be swing to at heat made by the contained moisture, or transferred from the apparent heat made by the contained moisture, or transferred from the apparent

y on constructed by means of the heard of the avena sensitiva, maind out.

ratus by its intervention, which checks the refrigeration; yet, ultimately, the whof the moisture being converted into vapour, the aggregate refrigeration does differ in the two cases.

225. Agreeably to Dalton's tables, at 70° the quantity of moisture in 31 grains 100 cubic inches of air, is 551-1000ths of a grain. The space allotted to this weight of vapour being doubled, it would remain uncondensed at 45° F., being associate with the same weight, but double the volume, of air; but at 32°, notwithstandithe doubling of the space, only 356-1000ths of a grain would remain in the aërife state; of course 551 — 356 = 195-1000ths, or nearly 2-10ths of a grain, would precipitated.

226. The latent heat given out by the condensation of this vapour, would heat, is well known, 1000 times its weight of water, or 195 grains, one degree; or grains 195-31ths = 6.29 degrees; and as the capacity of air for heat is only of fourth of that of water, it would heat 31 grains of air $6.29 \times 4 = 25.16$, or nea 25° F. As air, at 32° F., expands 1-480ths for each additional degree, the different of bulk, arising from the heat received, as above calculated, would be 25-480ths,

1-19ths nearly.

227. When air, replete with aqueous vapour, was admitted into a receiver p tially exhausted, and containing liquid water, a copious precipitation of moisture sued, and a rise of temperature greater than when perfectly dry air was allowed enter a vessel containing rarefied air in the same state. In the instance first m tioned, a portion of vapour arises into the place of that which is withdrawn dur the partial exhaustion. Hence when the air, containing its full proportion of pour, enters, there is an excess of vapour which must precipitate, causing a clo and an evolution of latent heat from the aqueous particles previously in the aerife state. As the enlargement of the space occupied by a sponge, allows, proportion bly, a larger quantity of any liquid to enter its cells, so any rarefaction of the when in contact with water, consequent on increase of heat or diminution of particles. sure, permits a proportionably larger volume of vapour to associate itself with given weight of the air. When, subsequently, by the afflux of wind replete w aqueous vapour, the density of the aggregate is increased, a portion of the vap equivalent to the condensation must be condensed, giving out latent heat, except so far as the heat thus evolved, being retained by the air, raises the dew point.

228. Hence, whenever a diminution of density of the air inland causes an interpolation of sea air to restore the equilibrium, there may result a condensation of aqueous pour, and evolution of heat, tending to promote an ascending current. This proposing followed by that which Mr. Espy has pointed out, of the transfer of heat fivapour to air, during its ascent to the region of the clouds, and consequent prectation of moisture, is probably among the efficient causes of those non-electrical storms, during which water from the Gulf of Mexico, or from the Atlantic, is tri

ferred to the soil of the United States.

Of the Influence of the Atmosphere in promoting Evapo. tion.

229. It has been seen that by its pressure the atm sphere opposes vaporization; yet a free access of air found indispensable in the desiccation of hay, or in t evaporation of water or other solvents. It was at c time generally conceived that evaporation resulted from affinity between the liquid and the air, analogous that between water and sugar, or alcohol and resin; t in consequence of the observations of several distinguish philosophers, a different view of this subject has been be terly entertained. It has in fact been ascertained that t quantity of aqueous vapour, in any space having sufficiences to liquid water, is always directly as the temper ture, whether there be a plenum or a vacuum, or whatever may be the density of the air simultaneously present.

putides promotes evapora: conty to which the va ur of clining itself through greater distance. a current of atmospheric only by removing the nel otherwise be exposed, ospheric interstices to a

231. Nevertheless, it appears to me that the influence a current of atmospheric air, in promoting evaporation, creater than can be reasonably thus accounted for.

232. It is difficult to conceive that the elements of atcospheric air should have no affinity for those of liquids; or that, if such affinity exist, it should not promote the woods of evaporation Nothing can be more certain that that eva poration is accelerated in proportion to the entet to which contact may induced between the aëriom and liquid particle! Ł e when surfaces, moistened lphuret of carbon, or the such volatile liqui s rolatile effers, are 1 to the wind, or to a blast, stance cold is produced by the accelerated evaporation. I is well known that the direction of the wind becomes widest from the sensation of coldness, experienced in that part of the wetted finger on which it blows. With the refigurating influence of a seze, when the skin is moistened by parapiration, we are all familiar.

the cense of ebullition, cannot be confounded in practice, however they may be identified agreeably to prevailing theories. In either case, heat is requisite, though much less in necessary in that of evaporation; but other things being equal, the process last mentioned, is accelerated in proportion to the extent of surface exposed to the air, while ebullition takes place in proportion to the surface exposed to the fire, without access of air. It only requires that the repour generated should have an aperture sufficient to allow of its escape, without increase of pressure. Hence the proportion of the surface exposed to the fire, without increase of pressure. Hence the proportion of the surface exposed to the fire, without increase of pressure. Hence the proportion of the surface exposed to the fire, without increase of pressure. Hence the proportion of the surface exposed to the air, while the fire without increase of pressure. Hence the proportion is accelerated in proportion to the surface exposed to the air, while the fire without access of air. It only requires that the proportion is accelerated in proportion to the surface exposed to the air, while the fire without access of air. It only requires that the proportion is accelerated in proportion to the surface exposed to the air, while the fire without access of air.

Call produced by the Evaporation of Ether when accelerated by a Current of Air.

The cold, produced by evaporation accelerated by air, may be advantageously shown by subjectir bulb simultaneously to a jet of ether,

and a blast from a bellows, so that the aërial and ethereal particles may be thoroughly mingled just before reaching the bulb. Water may be frozen in a bulb thus refrigerated.

235. Agreeably to the principle above illustrated, (217) that when air is liberated from a state of compression, cold ensues, I have lately contrived a new mode of exhibiting the vaporization of ether, so as to freeze water on a more extensive scale, and on a much more striking manner than heretofore. Between the lower part of a very strong vessel of sheet iron, capable of holding 40 gallons, and the "hydrent" pipes by which our city is supplied with water, a communication is made by means of a pipe and cock, so as to be opened or closed at pleasure. The vessel is previously filled with air, by allowing it to discharge any water which it may hold through a cock. Under these circumstances, on opening the communication with the hydrant pipes, the air within the vessel may be subjected to a pressure of more than one atmosphere. (154.) If by means of a suitable leaden pipe, furnished with a cock, and terminating with a capillary orifice, the air be allowed to blow into some ether and water contained in a thin capsule, the ether will be rapidly vaporized, and the water soon frozen.

236. In this experiment, in lieu of hydric (sulphuric) ether, we employ the new form of hyponitrous ether which I have lately discovered, the congelation will be

more rapidly accomplished.

237. It will hereafter be shown, that, by analogous causes, when solid carbonic acid is thrown into ether, a refrigeration is produced by which mercury may be rapidly frozen.

Definition of Vapour by Berzelius.

238. Berzelius objects to the use of the word vapour as implying a condensible acriform fluid. He uses it in the sense in which English authors employ the word fog, or cloud. Vapour and steam were originally, and still are used in this sense, yet the fluid which is used to propel steam engines, and to which they owe their distinguishing name, can only consist of water in the acriform state in which it is by the distinguished Swede designated as aqueous gas. Johnson defines steam to be the smoke or vapour of any thing hot and moist. Of course steam smoke and vapour have in some cases been used synonymously. I have elsewhere mentioned that before Black's discoveries and inferences were published, atmospheric air was the only acriform fluid whose existence was recognised. Hence the use of the words steam and vapour has grown with our knowledge, and consequently the names applied to visible steam or vapour have been extended to mean the invisible acriform fluids from which it is produced by refrigeration. I have some repugnance to designating by a common epithet, permanent gases, and the condensible clastic fluids produced from liquids above their boiling points. I do not see that any disadvantage arises from the customary use of the word vapour to designate the latter.

Of the Opponent Influence of Pressure on the Extrication of Gaseous Substances from a state of Combination.

239. When one of the ingredients of a solid or liquid is prone to assume the aëriform state, its extrication will be more or less easily effected, in proportion as the pressure of the air is diminished or increased.

prof Carbonie Acid from ! proacted by Echausie

of Lime subjected to an Acid, aked by Condensation.



240. If a tall cylindrical jar, containing a carbonate undergoing the action of an acid, he placed under a receiver, and the air withdrawn by an air comp, the effervencence will be augmented. But if, on the other hand, the same mixture be placed under a receiver, in which the pressure is increased by condensation, the effervescence will be dimimished. In the one case, the effort of the carbonic acid to assume the gaseous state is repressed; in the other, facilitated. Hence the advantage of condensation in the process for manufacturing carbonic acid water. Beyond an absorption of its own balk of the gas, the affinity of the water is imadequate to subdue the tendency of the acid to the aëriform state; but when, by mechanical pressure, a great number of volumes of the gas are condensed into the space ordinarily occupied by one, the water combines with as large a volume of the condensed gas, as if there had been no condemention.

inproved Apparatus for storing the influence of Pressure on Efferenceses.

Wh. A cylindrical speciwer, about 30 inches in height, and 3 inches in diameter, is appared on a westen block, W, between upright iron rods, RR, each at the least and, riveted to a plate of iron beneath the block, and, at the upper end, a sew at and furnished with a nut. By means of these screws and nuts thus formed, and anutervening evens bar, B, a brass disk, D, is pressed upon the rim of the retire. The disk is seground to fit to the rim of the glass, as that, with the aid of the best was duly seltened by lard, an air-tight juncture may be made. In the middle the disk there is an aperture, from which proceeds a stout tabe, with a mater each eide, severally furnished with gallows acrews, by means of which had pipe may be made to communicate with an air pump on one side, and a condense on the other. The tube is also surmounted by a cock, into which a glass functionmented. Before closing the receiver, some solid carbonate in pieces must be material as a security about one-third of the cavity. For this purpose I have material are one as to eccupy about one-third of the cavity. For this purpose I have material are as a transmitted of these substances, carbonic acid and lime are the principal ingression. In either of these substances, carbonic acid and lime are the principal ingression. The means being introduced, and the disk fastened into its place, as responsed in the figure, diluted muriatic acid may be added, by means of the funnel test each, in quantity enflicient to cover the carbonate.

The examination to cover the carbonate.

The companies of the superior affinity of chlorine for the calcium, and of hydran for the oxygen, (in the oxide of calcium or lime) the carbonic acid is expelled him gasters form, causing a perceptible effervescence or fearing of the liquid.

The these circumstances, by means of the air pump, the atmospheric pressure thin the remiver be lessened, the effervescence increases strikingly. On the other had, s, by closing the communication with the air pump, and opening that with the undersor while this is in operation, the pressure he increased, it will be seen the effervencence is diminished proportionably.

The experiment is much facilitated by the employment of an air nump.

The experiment is much facilitated by the employment of an air pump, with I have contrived, by which we can either exhaust or condense at pleasure. It Agreeably to experiments performed by Faraday, when the reaction between the still und a carbonate is made to take place in a stout tube hermetically scaled, the side may be separated in the liquid form. According to the more recent observation of the manufacture, this result has been attained upon a large scale, and one portion of the manufacture liquid has been found to be partially frozen, by the caloric abservation of the other portion.

245. Thilorier's process, as improved by Mitchell and others, will be hereafter il-474 lustrated and explained.

246. By analogous means various substances, naturally gaseous, have been liquefied

by Faraday, as will be mentioned in treating of those substances.

247. All the cases of liquefaction alluded to, are referable to the law that the power of any matter to pass to the acriform state is, ceteris paribus, less in proportion as the pressure is greater.

Of the Screw Rod and Plate Frame, employed in the preceding and many other Experiments.

248. The means by which the glass receiver, employed in the preceding experiment, is upheld and rendered air-tight by the rods, R R, the wooden block, W, the bar, B, and circular plate or disk. D is one to which I shall recent framently in the bar, B, and circular plate or disk, D, is one to which I shall resort frequently in the course of my experiments. Hence, to avoid unnecessary recurrence to analogous description, I shall in future designate as a screw rod and plate frame, that portion of the apparatus above described, which consists of the block, bar, plate, and screen rods.*

249. The glass in this case is made quite true by grinding on a large lap wheel, such as is employed by lapidaries. The same object is effected in the case of brass plates without grinding, by turning them in a lathe with a slide rest, and by a tool

with a fine pyramidal point.

OF CAPACITIES FOR HEAT, OR SPECIFIC HEAT.

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250. The power of equal weights of different substances, at the same temperature, in cooling or warming a liquid at a temperature different from their own, will be found very unequal. Thus the effect of a given weight of water being 1000, the effect of the like weight of glass will be 137, of copper 94, tin 51, lead 29, iron 110, gold 29, platinum 31, zinc 92, silver 55. If equal weights of water and mercury, at different temperatures, be mixed, the effect on the water will be no greater than if, instead of the mercury, 18th of its weight of water, at the same temperature as the mercury, had been added; and it takes twice as much mercury by measure as of water heated to the same point to have the same influence.

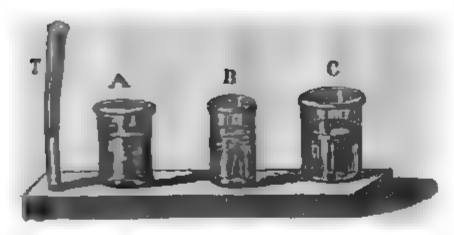
251. The term specific heat is usually employed to designate the quantity of caloric in a body in proportion to its weight or bulk, as specific gravity is used to convey an

idea of weight compared with bulk.

252. In the process above described, the specific heats of substances are found in order to estimate their capacities; the one being necessarily as the other, and the same series of numbers expressive of either.

[&]quot; Modification of the screw rod and plate frame are represented in the engraving referred to page 28. (153.)

Apparatus for illustrating Capacities for Heat.



363. Let be vesseles A., IB, and C, be : with 1 through the is, I, while communication with stal channel OL through block. The water was in all. state restance made by the : m ď med this liquid will be the our , : œ wilı common of water in the tube, 1. time total u une wande the index of the quant y þу ut u đ simpresion that they had all reno that the quantities sevi diffe A LOCOTA r huizonal aron. Of cours t ai the water within the vest a a fur s tabe, as any evidence of an eq 'n portions olby them.

Which measures the height of the mercury in the thermometer shows in maintance which substances, whose temperature it measures, are making my further accession of caloric; but it does not indicate the quantities, packedly received by them, in attaining the temperature in question is writes, in them, in proportion to their attraction for this self-repellent id; as the quantities of water received by the vessels, A, B, C, are varied the min of their respective areas.

Estimate.—It may be conjectured that this diversity in the power mbunaces, equally hot or cold, in influencing temperature, is due to a factor in their capacity to attract caloric, in consequence of which it which farms denser atmospheres about the atoms of some substances, as it does about those of others.

256. An analogy has already been suggested to exist between the mantrin which these calorific atmospheres surround atoms, and that in which searth is surrounded by the air; and also the mode has been suggested in high changes of temperature in the external medium would operate upon the density of such atmospheres. Supposing these preliminary suggesion correct, it would follow that the quantity of caloric absorbed or given that each exterior change of temperature, by any one congeries of atoms, while to that absorbed or given out by any other congeries, as the pretice condensation of caloric in the one, is to its previous condensation in the char. (173, 174, 175, 176.)*

Antice of the destrine of Petit and Dulong that the expecities of all elementary was fig heat are the same, will be deferred till I have treated of atomic pro-

Of the Specific Heat of Gaseous Bodies.

257. It was suggested by Lambert and Pictet, and the suggestion was afterwards sanctioned by Dalton, that space may have a capacity for caloric. Consistently with this idea the quantity of caloric in a given space should : always be the same whatever may be the gaseous fluid occupied by it. This is consistent with the fact that all the gases have the same capacity for heat, and all undergo a like expansion, in consequence of a like increase of Agreeably to this view of the case, the cold produced by rarefaction, as in the experiment with the exhausted receiver (212) or the palm glass, (215,) the heat consequent to the compression of air (219) arises from the caloric in the air or vapour, being too little for the space; allotted to the air in one case, and too great for that allotted in the other. This idea seems to have been abandoned in consequence of an experiment performed by Gay Lussac. This eminent chemist having made a Torricellian vacuum within a tall cylindrical glass receiver, about 3 inches in diameter and 39 in height, found that when the mercury employed was made to rise or sink in the vacant space so as alternately to enlarge or diminish it, no consequent variation of the temperature took place, since a delicate air thermometer, of which the bulb was included, indicated no It appeared, nevertheless, that when a minute quantity of air was admitted, any increase or diminution of the void space, consequent to the rise or fall of the mercury, was as productive, as the same thermometer showed, of a corresponding increase, or diminution, of sensible heat. Hence it has been inferred that a perfectly void space has no capacity for heat, the changes of temperature, consequent to the rarefaction or condensation of aëriform fluids, being altogether caused by corresponding changes in the capacity of those fluids for caloric. But as a perfect vacuum must liberate heat with perfect facility, it appears to me that the caloric should be absorbed by the mercury as rapidly as this metal could be made to encroach upon the space occupied by the calorific particles, and that, consequently, no palpable condensation of them could be effected by the above described process resorted to by Gay Lussac.

258. Admitting that, for equal weights, the specific heat of air is seven times as great as that of mercury, that of space being the same by the premises, there could not have been a capacity greater than that of about 200 grains of the metal, whereas a very small stratum of this metal, equal to one-fourth of an inch, would, in the apparatus employed, amount to more

than a pound.

259. The following experiments appear to me to be irreconcilable with the idea that the heat acquired by air entering a space does not arise from the specific heat of the space. When a receiver was exhausted so as to reduce the interior pressure to one-fourth of that of the atmosphere, and one-fourth was suddenly admitted, so as to lower the mercurial column in a gauge from about 22½ inches to 15 inches, heat was produced; and however the ratio of the entering air to the residual portion was varied, still there was a similar result.

260. When the cavity of the receiver was supplied with the vapour of ether, or with that of water, so as to form, according to the Daltonian hypothesis, a vacuum for the admitted air, still heat was produced by the latter, however small might be the quantity, or rapid the readmission. When the receiver was exhausted, until the tension was less than that of aqueous vapour at the existing temperature, so as to cause the water to boil, as in the Cryophorus, or Leslie's experiment, still the entrance of $\frac{6}{1000}$ of the quantity requisite to fill the receiver caused the thermometer to rise a tenth

of the cock, through one-fourth Crespon An of a circle, within one-third of a or time, was adequate to produce 1

the dange last mentioned.

38. The fact, that heat is produced, when to air, rarefied to one-fourth of the strespheric density, another fourth is added, seems to me to be irresachible with the idea, that this result arises from the compression of the parties of air previously occupying the cavity, since the entering air that he as much expanded as the residual portion is condensed.

362. As, agreeably to Dalton, a cavity occupied by a vapour acts as a needs to may air which may be introduced, I infer that when a receiver, the being supplied with ether or water, is exhausted so as to remove all the air, and lease mothing besides aqueous or ethereal vapour, the heat, acpixed by air admitted, cannot be ascribed, consistently, to the condensa-

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in of the repour.

368. It was ascertained by De of a thresometer is subjected to a mains, that the instrument sh he it be allowed to continue, a **ishmi by them, that** in the fir neut absorption of caloric; in a consequence of the c ment of its influx, had been l is my experiments above o the tes rapid, and the quantity or t tole religionated by rarefaction in auch condensed as to become

re and Marcet, that when the bulb of air while entering an exhausted refrigeration takes place. But if perature ensues. Hence it was efrigeration, and a con-1y an evolution of this prinm or air, which at the first s to me, nevertheless, L It a upon the thermometer ng air too minute, to allow it , and yet afterwards to be m by the evolution of caloric.

OF THE SLOW COMMUNICATION OF HEAT, COMPRISING THE CONDUCTING PROCESS AND CIRCULATION.

Of the Conducting Process in Solids.

364. It is well known that if one end of a piece of metallic wire, as a common pin for instance, be held in a cardle flame, the other end soon becomes too hot for the fagers. It is also known that the heated irons, used in soldering and other processes in the arts, have usually wooden handles, which do not become unpleasantly warm, when the irons within them are hot enough to blister the This inferior power of wood in conducting heat is also well exemplified by the handles of silver tea-pots, which are sometimes altogether of wood; in other instances principally of metal, small portions of wood inter-In either case, the facility with which the heat is spagated in the comparatively thin metallic socket, is contrasted with the difficulty which it experiences menting the wood.

An indicating ity of conducting power, when coms, is also displayed by common bone,

blue, iv y, porcelain, and especially glass.

Inequality of Conducting Power, experimentally illustrated.



of metal, wood, glass, and whalebone, each rod, at the end which is not cement to the wax, be successively exposed to the flame excited by a blow pipe. It will found, that the metal becomes quickly be ed throughout, so as to fall off from the wax; but the wood or whalebone may destroyed, and the glass bent by the ight tion, very near to the wax, without metal it so as to liberate them.

Additional Illustration.

267. The following method of illustrate ing the diversity of conducting power, pressed by different substances, has be suggested by an analogous process described in Silliman's Chemistry.

268. Rods similar in diameter and length, and consisting severally lead, tin, iron, copper, wood, and ivory, are made to pass from side to si through a vessel of sheet copper, in the shape of an oblong parallelopip Each rod extends on one of the sides, to an equal distance beyond the v sel. By these means, when the vessel is filled with boiling water, equiportions of each rod being situated within the boiler, they are all exposed an equal degree of heat. It is presumed that under these circumstances conducting power will be nearly in the inverse ratio of the time necesse to communicate to the equidistant ends of the rods, a heat adequate to can the ignition of similar pieces of phosphorus, simultaneously placed up them, before the application of the boiling water.

Rationale of the Fracture of Glass or Porcelain by Heat.

269. The fracture of glass or porcelain, exposed to fire, is the conquence of an inferior conducting power; as the heat is not distributed w quickness enough to produce a uniform expansion. Hence glass is as lial to crack by heat, in proportion as it is thinner. It may be divided by heated iron, by a string steeped in oil of turpentine and inflamed, or by theat generated by friction. (322, &c.)

Of the Conducting Power of various Metals.

270. Metals are by far the best conductors of caloric. There are, ho ever, scarcely two that conduct it equally well.

271. Despretz has ascertained by exact experiments, that the conduing power of the following metals is in the ratio of the subjoined number

4		•	,					•		
Gold,	•	•	•	-	•	•	•	•	1000.0	
Silver,	•	•	•	•	-	-	-	•	973.0	
Copper,	•	•	•	•	•	•	•	•	898.0	
Platinum,	•	-	•	-	•	•	•	•	381.0	
Iron,	•	•	•	•	•	•	•	-	874.3	
Zinc.	•	•	•	•	•	•	•	•	368.0	

49

CALORIC.

Tat. - - - 303.9 Lead, - - - 179.6

Explanation of the Process by which Heat is supposed to be communicated in Solids.

272. I conceive that in solids, the stratum of atoms forming the surface first exposed to the heat, combining with an excess of this principle, divides it with the next stratum. The caloric received by the second stratum, is in the next place divided between the second and third stratum. In the mean true the first stratum has received an additional supply of caloric, which process to the second and third stratum as in the first instance; while the quantity, at first received by them, is penetrating further into the mass.

273. It is I trust easy to conceive that, by the process thus suggested, called may find its way throughout any body, for the particles of which it may have sufficient affinity. Probably the superior conducting power of netals is due in great measure to a proportionably energetic affinity for calcure.

774. The conjectures, which I ventured to advance respecting the mode a which caloric may exist in atmospheres about atoms, seem to be pecuiarly applicable to the case of metals, on account of their great expansitility by heat, and susceptibility of contraction by cold. (174.)

275. If caloric be not interposed in a dense repulsive atmosphere between walks atoms, how can its removal cause that approximation of those atoms wards each other, without which the diminution of bulk invariably consequent to refrigeration could not ensue?

Liquids almost destitute of Conducting Power.

276. That liquids are almost devoid of power to conduct heat, is proved by the inflammation of ether over the bulb of an air thermometer, protected only by a thin stratum of water.

277. The inflammation of other upon the surface of water, as represented in this figure, does not cause any movement in the liquid included in the bore of the air thermometer at L, although the bulb is within a quarter of an inch of the flame. Yet the thermometer may be so sensitive, that touching the bulb, while under water, with the fingers, may cause a very perceptible indication of increased temperature. By placing the sliding index, I, directly opposite the end of the column of liquid in the stem of the thermometer, before the other is inflamed, it may be accurately discovered whether the heat of the flame causes any movement in it.



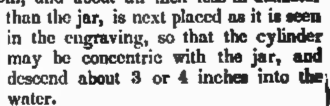
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Communication of Caloric by Circulation.

278. That caloric cannot be communicated in liquids, unless it be so applied as to cause a circulation of the particles, is demonstrated by the

following experiment.

279. A glass jar, about 30 inches in height, is supplied with as much water as will rise in it within a few inches of the brim. By means of a tube descending to the bottom, a small quantity of blue colouring matter is introduced below the colourless water so as to form a stratum as represented at A, in the engraving. A stratum, differently coloured, is formed in the upper part of the vessel, as represented at B. A tin cap, supporting a hollow tin cylinder, closed at bottom, and about an inch less in diameter



280. The apparatus being thus prepared, if an iron heater, H, while redhot, be placed within the tin cylinder, a the coloured water, about it, soon boils; 🛬 yet neither of the coloured strata intermingles with the intermediate colourless mass; and on sliding the finger upwards, while in contact with the glass, the heat will be found to have penetrated only a very small distance below the tin cylinder. But if the ring, R, be placed, while red-hot, upon the iron stand which surrounds the jar at 88, the portion of the liquid coloured bine, being opposite to the ring, will rise until it encounters the warmer, and of course lighter, particles, which have been in contact with the tin cylinder. Here its progress upwards is arrested; and, in consequence of the diversity of the colours, a well defined line of separation becomes conspicuous.

281. The phenomena of this interesting experiment may be thus explained.

282. If the upper portion of a vessel, containing a fluid, he heated exclusively, the neighbouring particles of the fluid being rendered lighter by expan-



sion, are more indisposed, than before, to descend from their position. But if the particles, forming the inferior strata of the fluid in the same vessel, be rendered warmer than those above them, their consequent expansion and

twice of aperia, which not have an artiful are in principally to the lower part of twice of temperature in a continue of temperature in a continue of temperature in the case or in the case of temperature

to give place to particles above ier. Hence heat must be apin order to occasion a uni-

w er i be aëriform or form mfluence of presco-oj and at others op-

wimmed Mostration of the Process by which Caloric is distributed in a Liquid until it boils.



284. On the first application of heat to the bottom of a vessel containing cold water, the particles in contact with the bottom are heated and expanded, and consequently become lighter than those above them. They rise therefore, giving an opportunity to other particles to be heated and to rise in their turn. The particles which were first heated, are soon comparatively colder than those by which they were displaced, and, descending to their primitive situation, are again made to rise by additional heat and enlargement of their bulk. Thus the temperatures reversing the situations, and the situations the temperatures, an incessant circulation is maintained, so long as any one portion of the liquid is cooler than another, or in other words, till ebul-

takes place; previously to which every particle must have combined as much caloric as it can receive, without being converted into steam. 55. The manner in which caloric is distributed throughout liquids by setting, as above described, is illustrated advantageously by an experit contrived by Rumford, who first gave to the process the attention in it deserves.

Into a glass nearly full of water, as represented by the foregoing a, small pieces of amber are introduced, which are in specific gravity tariy equal to water, as to be little influenced by gravitation. The most part of the vessel being subjected to heat while thus prepared, increased amber are seen rising vertically in its axis, and after they have surface of the liquid, moving towards the sides, where the vessel have from the influence of the external air. Having reached the sides in vessel, they sink to the bottom, whence they are again made to rise thre. While one set of the pieces of amber are at the bottom of the ill, some are at the top, and others at intermediate situations; thus de-

The state of specific gravity of the liquid.

monstrating the movements by which an equalization of temperature is ac-

complished in liquids.

287. When the boiling point is almost attained, the particles being nearly of the same temperature, the circulation is retarded. Under these circumstances, the portions of liquid which are in contact with the heated surface of the boiler are converted into steam, before they can be succeeded by others; but the steam thus produced cannot rise far before it is condensed. Hence the vibration and singing sound which is at this time observed.

288. According to an observation of Gay-Lussac, water boils in metallic vessels at a temperature nearly two and a half degrees lower than in those of earthenware.

QUICK COMMUNICATION OF HEAT, OR RADIATION.

289. It must be evident that the heat which we receive from a fire in opposition to the draught, reaches us neither by the conducting process nor by circulation. Actual contact is evidently indispensable to the passage of heat in either of these modes. The aëriform matter which is in contact with the embers, or the blaze of a fire, forms part of a current which tends rapidly towards the flue, as must be evident from the celerity with which the sparks which accompany it are propelled. The rapidity with which the aërial particles, heated by the fire, are thus carried up the chimney, far exceeds that with which caloric can be communicated, in the opposite direction, either by the conducting process or by circulation.

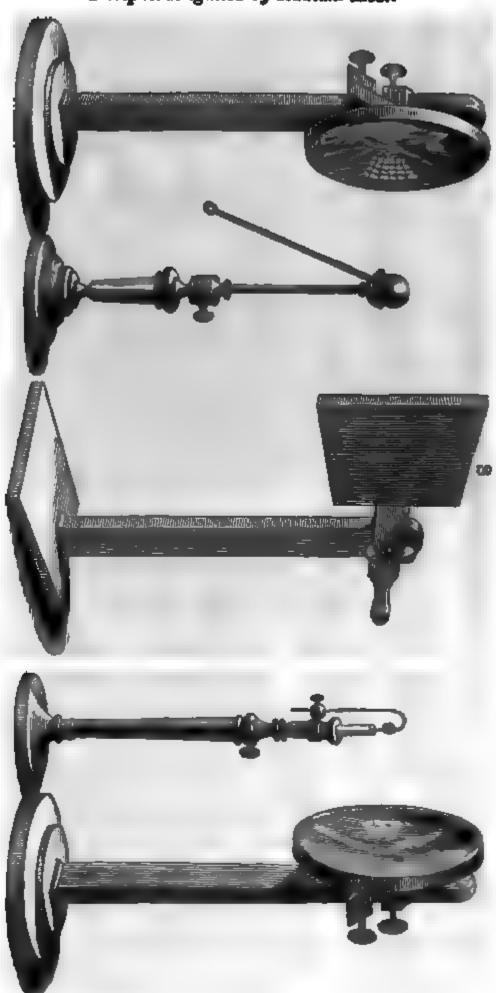
290. The caloric received from a fire under the circumstances above mentioned, and which is analogous to that by means of which the culinary operations of toasting and roasting are accomplished, is called radiant caloric or more usually, radiant heat. It has been called radiant because it appears to emanate in radii or rays from every hot or even warm body, as light emanates from luminous

bodies.

291. Radiant heat resembles light also in its susceptibility of being reflected by bright metallic surfaces; in which case it obeys the same laws as light, and is of course liable, in like manner, to be collected into a focus by concave mirrors.



Phosphorus ignited by Radiant Heat.



(Page 53.)

ted by this diagram, is to explain

in collecting the rays of radiant

pody in the focus of the mirror,

ig in intensity as the space into

a by the mirror, are reflected

Being thus corrected of their

it any other loss than such as arises

other mirror, which should be so

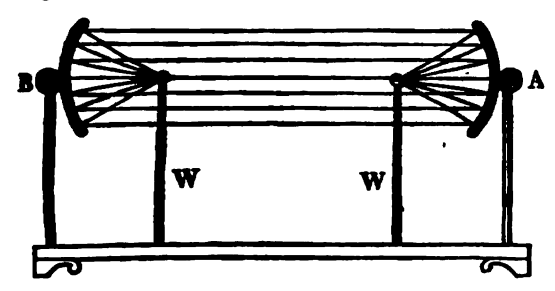
centres, from A to B, may at the

ented by the little balls supported

as the squares of the dis-

rating them in another.

Model for illustrating the Operation of Concave Mirrors.



32. The object of the model repre teme in which two mirrors open intented from one focus, and in conc

The caloric emitted by a h hould pass off in radii or rays, l hidthey pass enlarges; or, in o has But those mays which are an ter in directions parallel to its a: requery, they may be received, wi m acchanical imperfections, by had that the axis of the two m rrors may be coincident; or, in other wh, so that a line drawn through t time pass through their foci, 1 H be vires, W W.

The second mirror, B, reflects to its focus the rays which reach it on the first; for it is the property of a mirror, duly concave, to render this the divergent rays received from its focus, and to cause the parallel Which it intercepts to become convergent, so as to meet in its focus. The grings in the model are intended to represent the paths in

the rays move, whether divergent, parallel, or convergent.

Phophorus ignited at the distance of sixty feet by an incandescent Iron Ball.

26. The opposite engraving represents the mirrors which I employ in rightion of phosphorus and lighting a candle by an incandescent iron Linear produced this result at sixty feet, and it might be always efat that distance, were it not for the difficulty of adjusting the foci with ficient accuracy and expedition. I once ascertained that a mercurial twometer, when at the distance last mentioned, rose to 110 degrees of habeit

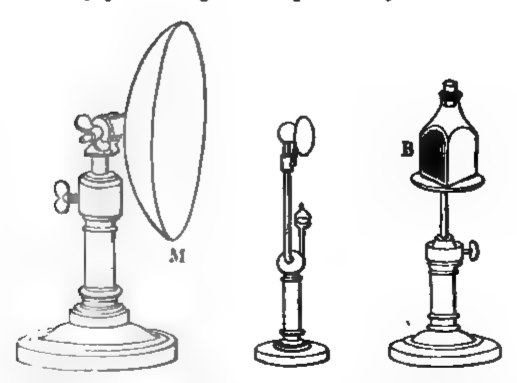
A tallow candle is so situated, that its wick, previously imbued with horus, may be in the focus of one of the mirrors. A lamp being rly situated with respect to the other mirror, it will be easy, by rethe focal image of the flame on any small screen, so to alter the This image to fall upon the phosphorus. This big ficed, the een, S, placed between the mirrors, is lowered so as

^{*} The single a mirror is in a line drawn from its centre through its true focus.

to intercept the rays. The iron ball being rendered white-hot is now sub-a-stituted for the lamp, and the screen being lifted, the phosphorus takes fire; and the candle is lighted.

Of the Diversity of Radiating Power in Metals, Wood, Charcoal, Glass, Pottery, &c.

Diversity of Radiating Power experimentally illustrated.



298. At M, (see figure,) a parabolic mirror is represented. At B is a square glass bottle, one side of which is covered with tin foil, and another so smoked by means of a lamp as to be covered with carbon. Between the bottle and mirror, and in the focus of the latter, there is a bulk of a differential thermometer, protected from receiving any rays directly from the bottle by a small metallic disk. The bottle being filled with boding water, it will be found that the temperature in the focus, as indicated by the thermometer, is greatest when the blackened surface is opposite to the mirror, and least when the tin fail is so situated; the effect of the naked glass being greater than the one, and less than the other.

299. The worst radiators are the best reflectors, and the best radiators are the worst reflectors; since the arrangement of particles which is favourable for radiation is unfavourable for reflection, and vice versa.

300. A polished brass andiron does not become hot when exposed from morning till night to a fire, so near that the hand placed on it is scorched intolerably in a few seconds. Fire places should be constructed of a form and materials to favour radiation: flues, of materials to favour the conducting process. To preserve heat in air or to refrigerate in water, vessels should be made of bright metal. In the latter case, the brightness is beneficial, only because the surface cannot be bright without being clean. If soiled, its communication with the liquid would be impeded.

301. Rationale.—Metals appear to consist of particles so united with each other, or with caloric, as to leave no pores through which radiant caloric can be projected. Hence the only portion of any metallic mass which can will are its roughly reduction in the output of the caloric can.

yield up its rays by radiation is the external stratum-

CALORIC. 55

In the other hand, from its porosity, and probably also from its not receive within its pores tenaciously as an ingredient in its composition in the posses but little obstruction to the passage of that subtile principle, when in the radiant form; and hence its particles may all be seemed, and engaged in radiating any excess of this principle with which the engaged in radiating any excess of this principle with which the engaged in radiating any excess of this principle with which the engaged in radiating any excess of this principle with which the engaged by any heated body, to the emanations from which the been exposed. We may account in like manner for the passage of that subtile power of earthenware and wood.

is the same reason that calorific rays cannot be projected from the metal, they cannot enter it when projected against it from the contrary, they are repelled with such force as to be reserved as any perceptible diminution of velocity. Hence the superior

ப்படர் metallic reflectors.

It wand seem as if the calorific particles which are condensed betwo was of the metal, repel any other particles of their own nature which
maintaite towards the metallic superficies, before actual contact ensues;
where a account of mechanical imperfection, easily discernible with
the second and account of mechanical imperfection, easily discernible with
the second and account of mechanical imperfection, easily discernible with
the second account of mechanical imperfections as they are
the second account of mechanical imperfections.
Their influence, in this respect,
the second account of the excellence of their general contour, and is not prothe second account of mechanical imperfections.

Radiation of Cold.

A thermometer placed in the focus of a mirror indicates a decline of the rature, in consequence of a mass of ice or snow being placed before at the situation occupied by the bottle in the preceding figure. This can be structured has been considered as demonstrating the radiation, and the materiality of cold. For since the transfer of heat the rature has been adduced as a proof of the existence of a material rate field. It is alieged that the transmission of cold by the same property with the admitted as equally good evidence of a material cause of cold.

36. The fall wing is the explanation which I give of this phenomenon,

ages are to the opinion that cold is diminished heat.

In the particles that caloric exists throughout the sublunary creation, as an the particle for and partly free. The particles of the free caloric I support to the earth by the general attraction of all the matter in the particles of the free caloric I support to the partly free. The particles of the free caloric I support to exert a self-repellent power, which increases with its desiry, as a the case of aëriform fluids. The repulsive power of caloric leafures the particles; in the quantity, it follows that either a diminution or interpretature in any spot must equally produce a movement in the particles; in the one case from the spot which sustains the case, in the other towards it.

Similar the surface of a mirror to be subjected to the influence of which a diminution of temperature has been produced, the rows of which particles between the mirror and the space will move into the The removal of one set of the calorific particles from the surface of the wifer, must make room for another set to flow into the situations the ward. The curvature of the surface of the mirror renders it more than the particles to succeed which lie in the direction of the focus.

Of the Observations and Apparatus of Melloni.

309. By means of a thermo-electric pile, and a galvanoscope or mulplier, of extreme delicacy, Melloni has lately ascertained some interesti properties of heat-producing rays, which serve to show a marked different and, at the same time, a great analogy between them and the rays of light

- 310. Let there be provided three transparent plates, severally of alurock salt, and rock crystal or glass, each about an eighth or tenth of inch thick; it will be found that the effect of the transmitted rays upon to pile, when unimpeded, being 30, that which takes place during the interposition of the rock salt, will be 28, during the interposition of the rock salt, while during the interposition of the alum the effect would be two or three.
- 311. The effect of interposing a plate of smoky rock crystal, will, und the same circumstances, be equal to 14 or 15.
- 312. In other words, out of 30 parts, rock salt intercepts two parts the influence of the radiant heat; rock crystal, whether smoky or clear, tercepts about half; while alum, or glass, intercepts nearly the whole.
- 313. If, in like manner, two pairs of plates be employed, one pair form of a pane of green glass (impermeable to red rays,) and a plate of alu the other pair formed of a pane of perfectly opake black glass, coup with a plate of rock salt, it will be found that the first mentioned pair tercepts the calorific radiation entirely, while the other permits nearly of third as much to pass, as when not interposed.
- 314. Hence it appears, that bodies, quite permeable by light, may tirely intercept radiant heat, while others, impermeable by light, allow t passage of radiant heat. Melloni designates the former as athermane, t latter as diathermane bodies.
- 315. It follows that permeability to heat-producing rays is not to confounded with transparency.
- 316. Radiant heat has been found by Melloni to vary in its power permeating bodies, according to the source from which it proceeds, and t media through which it may have passed. After passing through nit acid, more will pass through alum than if received directly from the sour
- 317. Moreover certain media have, with respect to calorific rays, an fluence analogous to that which coloured media have with respect to lig in allowing some rays to pass, while others are arrested.
- 318. This property of the diathermane bodies, is called diathermans Rock salt seems to be a diathermane body, devoid of diathermansie. I last mentioned property lessens as the body is thinner, and may, as in a case of coloured media, be rendered null by an extreme tenuity.
- 319. The non-luminous calorific rays have been ascertained by Mello to be susceptible of refractions analogous to those of light. When thermo-electric pile is so situated as that the rays of heat cannot directeach it, by interposing a prism of rock salt, having a refracting angle 60°, the rays will be made to reach the pile.
- 320. From experiments performed by Prof. Forbes, of Edinburg, we the aid of Melloni's thermoscope, above alluded to, it appears that radia heat, unaccompanied by light-producing rays, is susceptible of polarization. Respecting this fact, some further mention will be made in treation of the polarization of light.

57

MEANS OF PRODUCING HEAT, OR RENDERING CALORIC SENSIBLE.

Of the Solar Rays as a Source of Heat.

21. Of all the natural sources of heat, the sun is ob-

vicesly the most prolific.

322. The solar rays may be collected into a focus either by the refracting influence of glasses or the reflecting power of mirrors. They may be converged by reflection, a mode analogous to that illustrated in the case of radiant heat.

323. The glasses employed for concentrating light are called lenses from their shape, which is that of a double

corez lens.

24. As the intensity of the heat produced by the solar least is in proportion to the quantity of them which may be collected upon any given spot, there appears to be no init to the degree of heat producible by their concentration, excepting that arising from the difficulty of making leases sufficiently large and free from defect, or of associating mirrors sufficiently numerous and well arranged.

325. Until lately, scarcely any occurrence of antiquity appeared more unaccountable than the destruction of the Roman ships, which Archimedes is alleged to have accomplished. by concentrating upon them the rays of the sun. Nevertheless, of this wonderful feat, Buffon seems to have discovered the means. Having arranged a number of plane murors so as to concur in reflecting the solar image upon the same spot, he was enabled to fuse lead at a distance This contrivance resembles that which Arof 140 feet. chimedes employed, if we may judge from the accounts which have been given of the latter. Previously to the employment of pure oxygen gas, the hydro-oxygen blowpipe, and voltaic electricity, there was no known mode of malling the heat produced by large burning-glasses and Birrors.

Sensible Heat evolved by Electricity.

226. The power of lightning to produce ignition is displayed by the conflagration of ships and barns, in concepte of the ignition of cotton, hay, or other combustible. The power of the electric spark to ignite an inflam-

mable gaseous mixture is agreeably illustrated, by means of the apparatus described in the following article.

Application of an Electrophorus to the Ignition of Hydrogen Gas, generated in a Self regulating Reservoir.

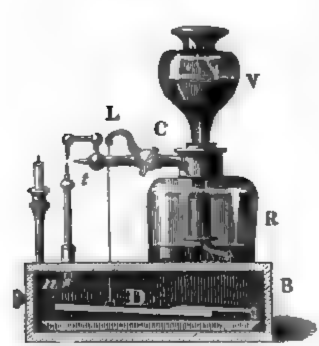
327. In order that the interior of this apparatus may be described, (see fig. be low) the side of the box, B, below the reservoir, B, is supposed to be removed. On the bottom of the box is a square metallic dish covered by a stratum of seeling was. The metallic plate, D, is supported behind by a glass rod, comented to a socke soldered to a hinge. Upon this hinge, like the lid of a trunk, the plate moves freely while connected with the lever, L, by a silken cord. The lever, L, is attached to the key of the cock, C; so that opening the cock causes the plate to rise, and touc the knob, a, of the insulated wire. This were terminates just before the orifice of the tube, t, proceeding from the cock, and about one-eighth of an inch from another wire, supported upon that tube.

wire, supported upon that tube.

328. The glass reservoir, R, receives into its open neck, the tapering part of glass vessel, V, which is so proportioned, and fitted to the neck by granding, as t

make with it an air-tight juncture.

329. Below this juncture, the vessel, V, converges, until it assumes the form of



tube, reaching nearly to the bottom to the reservoir. Around the tube the formed, a coil of gine is supported, a as to be above the orifice of the tube constituted as abovementioned.

supplied with diluted sulphuric action between this solvent are the reaction between this solvent are the rinc will evolve hydrogen gas. The gas thus evolved, if not allowed to a cape, will force the liquid which generates it through the orifice of all tube proceeding from the vessel, I into the cavity of this vessel, until the quantity of the acid remaining below is insufficient to reach the sine. When ever this takes place, the evolution a hydrogen ceases. As soon, however as, by opening the cock, any portion a the gas is allowed to escape, an equivalent bulk of acid descends into the reservoir, and reacts with the rise until, by the further generation a

hydrogen, the portion of acid which may have descended shall again be expelle from the lower into the upper vessel. At the same moment that, by turning th cock, C, a jet of gas is emitted, the plate of the electrophorus being lifted agains the knob, a, of the wire, an electrical spark will pass from the other end of this wire to that of the wire supported by the cock, and of course uninsulated by it communication with the operator's hand. Consequently the jet of bydrogen will be ignited, and will light a candle exposed to its influence.

331. For a rationale of the electrophorus, as also for other exemplifications of the igniting power of electric duscharges, I refer to my treatise on statical or mechanics

electricity.

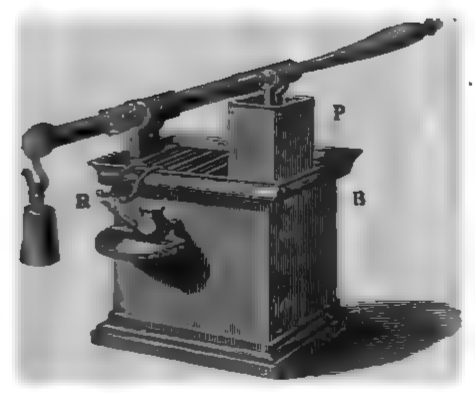
Ignition by Galvanum.

Galvanie Apparatus for Lighting a Lamp.

332. The following figure represents an instrument for lighting a lamp by mean of a galvanic discharge from a calerimeter; for a more ample explanation of which

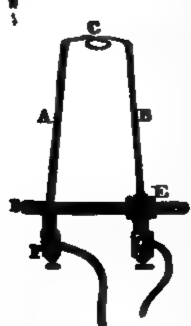
I must refer the reader to my lectures on galvanism.

333. The plunger, P, being depressed by means of the handle attached to it, some acid contained in the box. B, is displaced, so as to rise among the galvanic plate. By the consequent evolution of the galvanic fluid, a platinum wire, fastened between the brase rods forming the poles of the calorimotor, and projecting over the lamp accent at R, is rendered white bot, and a filament of the wick, previously laid upon a inflamed.



note as a counterpoise to the plunger, and when it is not depress-to it out of the acid.

Galtono ignition Apparatus.



235. In many of my experiments, for the purpose of producing the temperature of combustion in cavities inaccombine by ordinary means, I employ a wire ignited by being made a part of a galvanic circuit.

236. Of the apparatus by which this object is effected, I shall here give a description accompanied by a figure, which will convey a general idea of the contrivance, applicable to all cases where it may be used. Having thus recorded the student. I shall in fature re-Having thus propored the student, I shall in future refor to it under the name at the head of this article, in order to avoid sitemmlocution, and unnecessary recurrence to analogue description. D represents a section of a metallic disk. A B, two metallic rods, which should be of iron, if in contact with moreury, but which otherwise may be of bruse, are made to enter the cavity. If, as in general, the rode pass through a metallic plate or cylinder, one of them may be solder-ed to the plate or cylinder. The other must be so secured, where it passes through the metal, by a collar of leather, E, as to insulate it from all metallic con-

tact, and to render the aperture through which it en-ters, air-tight if necessary. The rods may extend into ty to any convenient distance, their terminations being approximated, more as may be desirable, but not brought in contact. To one of these rods, timetes within the cavity, one end of a fine platinum wire is coldered; d of the wire being soldered in like manner to the similarly situated tertibe other rod. To the rod secured by the collar of leather, at the termioutside of the cavity, a gallows acrow is attached, by means of which or copper red may be made fast at one end, while the other is fastened poles of a competent calorimotor. To the other pole of the calorimotor is attached at one end, which at the other may be secured by a galither seldered to the plate, or to the projecting extremity of the unin-in the figure. Sometimes the last mentioned rod is left at liberty, so "hen desirable, any part of the apparatus having a metallic uniconlated rod. If, under these circumstances, the cale-

han eposson, the wire will be ignited.

Ignition by Collision.

337. The ignition of spunk, tinder, or gunpowder, by means of flint and steel, comes under this head. In the rotary match box, the collision is produced by a wheel thrown into rapid rotation. An analogous apparatus, called a steel mill, had long been employed to procure light in mines infested with light carburetted hydrogen, prior to Sir H. Davy's invention of the safety lamp. This gas explodes on coming into contact with the flame of a lamp or candle, but is not ignited by the scintillations from a steel mill.

Heat produced by Percussion.

338. A rod of iron hammered with great rapidity by a skilful workman, will become so hot as to ignite a sulphur match, and phosphorus may be easily ignited in this way; but the same piece of iron cannot be ignited by percussion more than once.

339. Coins grow hot when struck in the coining press, but, if cooled during each interval between the blows, are less heated at each successive blow. At the same time the density of the mass is permanently increased, probably by the expulsion of the caloric, interposed between the metallic atoms. (272.)

Heat produced by Friction.

340. Friction, as a means of producing heat, differs from percussion; since in the case of friction, the effect being confined to the surfaces of bodies, there is no condensation of the mass subjected to the process. Collision differs both from percussion and friction; for it produces ignition only in the minute portions of matter which are struck off. The masses employed are not heated.

341. It is well known that savages avail themselves of the friction of wood to produce fire. Wood revolving in the lathe may be carbonized, throughout the circle of contact, by holding against it another piece properly sharpened. By rubbing one cork against another, sufficient heat is

produced to ignite phosphorus.

Glass so heated by the Friction of a Cord, as to separate into two parts on being subjected to Cold Water.

342. The process for dividing a tube, which I am about to describe, illustrates at

once the heat produced by friction, and the non-conducting power of glass.

343. Some years ago, Mr Isaiah Lukens showed me that a small phial or tube might be separated into two parts, if subjected to cold water, after having been heated by the friction of a cord made to circulate about it, by two persons alternately pulling in opposite directions. I was subsequently enabled to employ this process for dividing large vessels of four or five inches in diameter; and likewise to render it is every case more easy and certain, by means of a piece of plank forked like a boot jack, as represented in the following figure, and also having a kerf or slit cut by a saw, parallel to, and nearly equidistant from the principal surfaces of the plank, and at right angles to the incisions forming the fork.

344. By means of the fork, the glass is easily held steady by the hand of one operator. By means of the kerf, the string, while circulating about the glass, is confined to the part where the separation is desired. As soon as the cord smokes, the glass is plunged into water, or if too large to be easily immersed, the water must be thrown upon it. This method is always preferable when the glass vessel is so open, that on being immersed, the water can reach the inner surface. As plunging is the most effectual method of employing the water, I usually, in the case of a tube, closs the end which is to be sunk in the water, so as to restrict the refrigeration to the

outside.



Incience.—If the friction be continued long enough, the glass, though a very allocater of heat, becomes heated throughout in the part about which the friches plane; of course it is there expanded. While in this state, being suddenly sated by the cell water on the outside only, the stratum of particles immerally the cell water on the outside only, the stratum of particles immerally charge. Hence a separation usually follows: see (264, dec.)

Ignition by Attrition.

is by means of a lathe, the circumference be brought into contact with not steel, heat will be so copiously evolved at the place of collision, is steel may be actually divided by the successive ignition and abrata portion of its particles. The ignition is confined to the steel, better heat, evolved in this case, is too much divided upon the whole situace of the iron, to affect any part materially; whereas, a few sits of steel having to encounter successively many of iron, the heat, rated by the attrition, accumulates in the former, so as to produce visitation.

7. This case differs from that of pure collision, since, although heat is and in the abraded particles, it is also produced in the mass; and it shows that of friction, since, although both of the masses are heated, masset heat is evolved in the matter which is abraded.

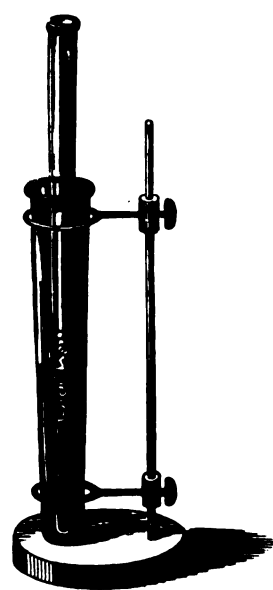
Heat produced by Combination.

The union of tin or lead with platinum is productive of a remarkinvation of temperature. For the exhibition of this phenomenon, both that he in the state of foil, and the more fusible metal rolled up in them, so as to form a scroll as large as can be conveniently ignited that blowpine. As soon as the scroll reaches a red heat, it betaken as a management of the union being effected with an as-

Experimental Illustration.

349. Tin foil and platinum foil are rolled up in scroll, the tin being innermost, and the whole subjeto the flame of the hydro-oxygen blowpipe, supplied currents of hydrogen gas and atmospheric air. Almos soon as the mass reddens, it becomes incandescent an energy almost explosive. (250, &c.)

Boiling Heat produced in Alcohol, by the Mixture of Sulphuric with Water.



350. The evolution of caloric, produce the mixture of liquids, has long been an of attention among chemists. A sensib crease of temperature arises from the mand consequent combination of alcohol water. When sulphuric acid is added to an analogous result ensues, but the rise of perature is much greater. The heat, thus rated, may be conveniently exhibited by a of the apparatus represented by the adjifigure, and the process which I am about scribe.

351. Into the inner tube introduce as alcohol, coloured to render it more discenses will occupy it to the height of three of inches. Next pour water into the outer till it reaches about one-third as high as quid contained in the inner tube; and after add to the water about three times its be concentrated sulphuric acid. The liquid inner tube will soon boil violently, so as t in a foam.

Solution the Means of producing Heat or Cold.

352. Solution produces either heat or cold, according to the nature substance dissolved and of the solvent employed.

353. In absorbing and dissolving gaseous ammonia or chlorohydrigas, the resulting liquid becomes hot. Water becomes cold in dissolving nitrate of ammonia. Sulphuric accomes at first boiling hot, and afterwards freezing cold by successive tions of snow.

Evolution of Caloric by Mechanical Action inducing Chemical D position.

354. With the view of showing the necessity of distinguishing heal latent cause from sensible heat, the explosion of a fulminating powd percussion was exhibited. This phenomenon falls under the defigiven at the head of this article. Ignition produced in this way has a been advantageously applied to fire-arms and fowling pieces. (30.)

caloric. 63

355. It seems probable that the mechanical force of the blow causes were redicted of the compound to be nearer to each other; in consequence of which an arrangement of the elements ensues, inconsistent with the remove of the large quantity of caloric with which they were previously consisted.

The inflammation of a friction match, appears to me to arise in part in meat generated by friction, and in part from mechanical impulse, interest a chemical reaction between the ingredients, and exposing them to mean the mean matches, which take fire when crushed, owe this result to the last medical cause only.

The rationale of the chemical reaction of the ingredients, will be

given under the heads of sulphur, phosphorus, and the chlorates.

Heat produced by Condensation experimentally illustrated.

35%. Spunk or tinder may be ignited, if introduced into a condenser of

expressive construction, and the air forcibly condensed upon it.

while during its condensation it becomes warmer. It seems that when the compression is carried very far, so much caloric is liberated as a came against This result is attained by means of a small condenser, the construction of which does not differ from that which has been described (145. &c.), excepting that a cock for the introduction of the spunk is substituted for the valves. The ignition is accomplished by having the piston attained, as that there may be as much air as possible included by it, and then driving it home, with a jerk, so as to condense the air upon the maner to be ignited with great force and rapidity. Sometimes the instrument is made of glass without a cock, so that the ignition may be seen; the stand with great into a cavity in the end of the piston, which must of the cold with application of the ignition is effected, in order to make any set of application of the ignited spunk.

It appears evidently from this phenomenon that, in air, the quantity is at the proportion to the ponderable matter lessens as the density in-Ur. nother words, as the space allotted to the air is diminished.

The inference would appear, at first view, irreconcilable with those is the which demonstrate that, in steam, the quantity of caloric is a part of as the weight of water; but the discordancy disappears with weight of that the heat of the condensed air is estimated after the same of the semble heat liberated by the compression; while in the case of them the came the cannot be permitted, as a loss of sensible heat would be attracted with a partial condensation, producing a proportionate diminution of description.

If spain, formed at the boiling point of 212°, and having no access kni-t. In the liquid form, were to be raised to some higher temperature, it might be subjected to compression without being parameter. It is that the same law would apply to it as to atmospheric which always exists at a heat far above its boiling point, and has no access that of its own kind of ponderable matter in the liquid form.

By the boiling point of air, I mean that temperature below which a ward become liquid. We have, I think, reason to infer that all aëriform for prove susceptible of liquefaction, if our ability to condense

* x x power of producing cold were unlimited.

364. It has been suggested (257, &c.), that the caloric thus condenses may belong to the space, and not to the air.

Experimental Illustration.

365. Spunk ignited in consequence of the compression of air, by means of an appropriate condenser.

Of Fermentation as a Source of Heat.

366. It is well known that vegetable substances, while undergoing fer mentation, acquire a great accession of heat; and that green hay is a times spontaneously ignited. The heat generated in stable litter is employed to sustain the temperature necessary to the corrosion of the metal in the manufacture of white lead.

Of Vitality as a Source of Heat.

367. The temperature of warm blooded animals demonstrates the power of animal life to evolve caloric. In no other respect is chemical reaction an analogous to that which takes place within the domain of vitality, as it their common association with heat, both as cause and effect. The old chemical law that bodies do not act unless fluid, to which the actual exceptions are but few, shows how much the processes of chemistry are dependent on the principle, without which there could be no fluidity. The dependency of life on temperature is self-evident. Seeds and eggs lie does mant until excited by a due degree of heat.

Of the Means of exciting or supporting Heat for the Purposes of Chemistry.

368. It is well known that the activity of fire is dependent on the supply of air, as well as on the quantity and quality of the fuel.

369. As the air which comes into contact with a fire is necessarily much rarefied by the expansive power of heat, it has consequently a tendency to ascend in a vertical current, giving place to the colder and heavier air in the vicinity, agreeably to the principles already illustrated. See (282 and (286). The limits of this vertical current of heated air, in the case of a smoky lamp flame, are well indicated by the fuliginous particles. It may however, be observed that the influx of the cold air takes place not only or a level with the flame, where it must quicken the combustion, but also above the flame, where it narrows the heated column and retards its progress. In the Argand lamp, a glass chimney defends the vertical current from lateral pressure, until it has attained a sufficient height to cause an adequate current of air to act upon the flame.

370. In conformity with the principle thus illustrated by this elegant and useful contrivance, all air furnaces are constructed. The hot air and vapour proceeding from the fire, being received into a flue, or the furnace being tall enough of itself to protect the ascending current, all the air which flows in to take its place is made to pass through the fuel.

371. It would not be expedient to take up the time of the student with a detailed explanation of the various furnaces used by chemists. Some of them will be introduced in subsequent illustrations, as associated with

processes, in which their utility and the method of using them will be

Experimental Illustration.

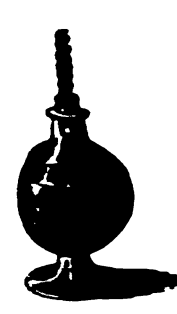
372. An Argand lamp shown and explained. Also an Argand lamp with concentric wicks.

Of the Bellows, and of Forge Fires.

373. The bellows is so universally known as the means of exciting common employed by smiths, as to render it scarcely necessary to mention the forge fire as among the most efficient and convenient methods of proteing heat for the purposes of chemistry. The supply of air is, in this case, yielded by an operation analogous to that of the condenser. (148, &c.)

In the double bellows, the additional compartment performs a part, in exalizing the efflux, equivalent to that of the air vessel in the case of the air pump, the valves operating in the same way. (143.)

Lamp without Flame.



375. About the wick of a spirit lamp, a fine wire of platinum is coiled, so as to leave a spiral interstice between the spiral formed by the wire; a few turns of which should rise above the wick. If after lighting a lamp thus constructed, the flame be extinguished by a gentle blast, or the transient application of an extinguisher, the wire will be found to remain red hot; as it retains sufficient heat to support the combustion of the alcoholic vapour, although the temperature is inadequate to produce inflammation.

Rationale.—The metallic coil appears to serve as a reservoir for the caissic, and gives to the combustion a stability, of which it would otherwise be deficient. There is some analogy between the operation of the way is setting as a reservoir of heat in this chemical process, and that of a fly wheel as a reservoir of momentum in equalizing the motion of machinery.

Of the Mouth Blowpipe.



As a fire is quickened by a blast from a bellows, so a flame may be excited by a stream of air propelled through it from the blowpipe. The mean known by this name, is here represented in one of its best forms. It is exceptible of various other constructions; all that is essential being a fixed a size at one end suitable to be received into the mouth, and to the other end having a bend nearly rectangular, beyond which the conserges to a perforation, rather too small for the admission of a

common pin. There is usually, however, an enlargement, as represe in this figure, to collect the condensed moisture of the breath.

378. The mouth blowpipe is of great service in assaying minute tions of matter, so as to form a general idea of their nature. The brated Berzelius, who has written an octavo volume on the subject of the strument, informs us that by means of it Gahn discovered tin in a min in which it had not been detected by analysis, although existing only it proportion of one per cent: also that he had often seen him extract a bule of metallic copper from the ashes of a quarter of a sheet of particles. The utility of the mouth blowpipe will be manifested in several future, trations.

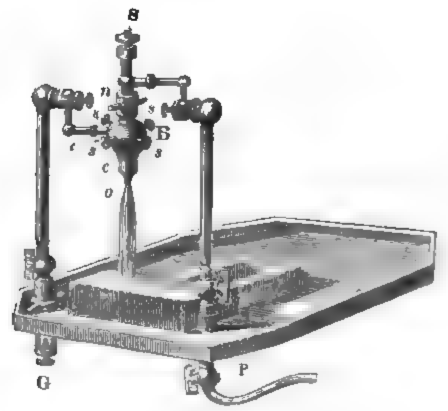
Of the Enameller's Lamp.

379. A lamp, so made as to be excited by a jet of air from a station blowpipe, supplied by a double bellows, gasometer or gas-holder, is empled much by chemists and artists for bending glass tubes, or heating them as to blow, on them, bulbs for thermometers. Such lamps having been ginally used by enamellers, are designated accordingly.

Of the Hydro-Oxygen or Compound Blowpips.

380 In the year 1801, by the invention of the hydro-oxygen or compound pipe, of which I published an account the following year, I was enabled to fur veral of the pure earths which had previously been deemed infusible; and like not only to fuse, but to volatilize pure platinum. Subsequently, my friend Prof Silliman, by a more extended use of the instrument, fused a great number of stances insusceptible of fusion by the common blowpipe. My memoir was us lished in London, in Tilloch's Magazine; also at Paris, in the Annales de Chand was noticed by Murray in his treatise of chemistry, and by Dr. Hope, it lectures; yet, when a modification of the hydro-oxygen blowpipe was contrive Mr. Brooke, Dr Clarke, by means of this modification, repeated my experis and those of Professor Silliman, without any other notice of our pretensions such as were calculated to convey erroneous impressions.

Engraving and Description of an improved Compound Blowpipe and its Append



381. The following figure represents a compound blowpape which I contrived executed myself in the year 1813; but, fearing it might be deemed unnecess

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I did not then publish an account of it. Experience has shown that the en of its structure does not render it more difficult to use than the simnaments intended for the same purpose; while its parts are popularly suc-

description adjustment.

s a brace ball, with a vertical perforation, terminating in a male screw in a formule acrew below. Another perforation, at right angles to this, magnanication with the tube t, which enters the ball at right angles. smaller breas ball may be observed above, with perforations similar to s larger ball, and a tube, in like manner, entering it laterally. This ball is a made screw below as well as above. The thread of the lower screw to the left, while that of the screw of the larger ball, which enters the n, in curved to the right. Hence the same motion causes the male screws h, or recode from each other, and thus determines the degree of compresto a cock which is placed between them in the nut. At S, above the ball, row may be observed, with a milled head. This is connected with a small h manu through the cork in the nut, n, and reaches nearly to the external hem which the flame is represented as proceeding. This tube is for the of base, but at its lower end terminates in a tube of platinum. It comby lateral apertures with the cavity of the upper ball, but is prevented by our communicating with the cavity in the other ball. Hence it receives nich may be delivered into the upper ball from the lateral pipe which enall, but receives more of the gas which may enter the lower ball, B.

• the formale screw of the latter, a perforated cylinder of brass, c, with a ing male acres, is fitted. The perforation in this cylinder forms a contithat in the ball, but narrows below, and ends in a small hollow cylinder

m, which forms the external orifice of the blowpipe, c. regit it from the cavity of the smaller ball. The intermediate nut, by ig about the tube the cerk which surrounds it, prevents any communicaon the cavities in the two balls. By the screw, 8, in the vertex, the oricoming trade may be adjusted to a proper distance from the external oriso different splinders, and as many central tubes with platinum orifices of alibers, were provided, so that the flame might be varied in size, agreeably et in view.

nve always deemed it best to transmit the oxygen gas through the tube in ince two velumes of the hydrogen being required for one volume of oxyarger tube eaght to be used for the former; and the jet of hydrogen is

:ween a jet of oxygen within it, and the atmospheric air without.

ider the table is a gallows, G, with a screw for attaching a pipe, leading

f-regulating reservoir of hydrogen.

order to put this apparatus into operation, it is affixed to a table, as reprethe figure, or to a smaller stand, and secured to the side of the hydror cistern, so as to be conveniently situated for receiving the oxygen from a z, through the pipe, P, and the hydrogen through a pipe attached at G. auther pipe, proceeding from a reservoir of hydrogen gas, is attached, by The screw and gallows, G, to one of the tubes communicating with the

be eavity of the hydrostatic blowpipe may be supplied, either with oxygen, phone air. In either case, in order to have the instrument in full operation,

measury to open the cocks duly, and inflame the hydrogen.

he heat preduced, in this way, by the combustion of hydrogen with atmosr, is sufficient to fuse platinum; and when oxygen gas is employed, that may other, may be volatilized. The facility with which the hydro-oxygen wither excited by pure oxygen or common air merely, may be made to act, averagent direction, renders it peculiarly serviceable in many operations; its issaliness is a great recommendation.

mond's Lime Light, and of Daniell's and Maugham's Bloropipe, so called erroneously.

has been said in some of the British newspapers, of the application in the light reflected by lime, when subjected to the flame of the com-This is treated as a new invention, although in my original Mein the year 1802, I spoke of the light so created as intolerable to the A similar observation will be found in the description given by my of the phenomenon in question. It follows that the ber can cal, ay claim to a new application of a previous discovery.

392. In my original memoir on the hydro-oxygen blowpipe, I described and represented by engravings two methods of causing the currents of the two gases employe to meet. Agreeably to one of these, two perforations were made to unite and form on at about the tenth of an inch from the external orifice, so as that a section of the a gregate would resemble in shape the letters XY. Agreeably to the other method, smaller tube was made to enter and to be concentric with a larger one, the latt being a little longer, so as that at a little distance from its end, the orifice of the former terminated. The oxygen being supplied through the inner tube, and the hadrogen through the outer one, the admixture of the oxygen with the hydrogen, to place within the bore of the external tube, at a small distance from its orifice.

393. Not being enabled to procure any platina at the time, I could not construct blowpipe, of the last mentioned kind, sufficiently refractory; but about the ye 1815, I constructed the compound blowpipe above described, and exhibited it to Pr fessor Silliman, who mentioned this fact in a letter written within a year afterware From the time that I was elected Professor of Chemistry in 1818, I have employed this form of the instrument, of which an engraving and description was given in t Franklin Journal (Vol. I, 1826, p. 195,) of a simpler instrument upon the same pri ciple, an engraving and description of which will be found in Silliman's Journal 1 1822. Yet both Professor Daniell and Mr. Maugham, resorted to analogous co trivances. The former has been called Daniell's hydro-oxygen blowpipe, the oth also is distinguished by the name of its contriver. It differs from mine essential only in being recurved into an acute angle, so as to throw the flame on a cylinder lime, for the purpose of illumination. In order to accomplish the same object, I has only to direct mine obliquely upwards, instead of resorting to a direction deviating little from the perpendicular, as is usually preferable. It is surprising that und these circumstances, Maugham should have received a premium for the instrume which he had thus modified, without any reference to the original inventor.

Improved Process for the Fusion of Platinum.

394. Latterly by multiplying the jets, and using great pressure, I have been en bled to fuse more than two pounds troy, of platinum, into a malleable mass. To method which I employed, was the same essentially as that described in Silliman Journal, as abovementioned. The gases are made to mingle in a common cavity and afterwards to supply jet tubes of about the usual size of those employed a blowpipes; these are to be made more or less numerous, in proportion to the quantity of metal to be fused. The great desideratum is to have the pressure on the gase sufficiently great, and at the same time perfectly steady.

MEANS OF PRODUCING COLD, OR RENDERING CALORIC LATENT

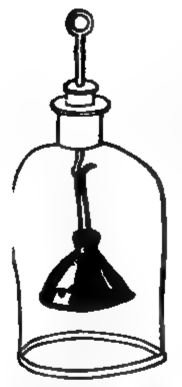
Cold by Vaporization.

395. The cold produced by evaporation has been illus trated by an experiment in which a jet of other, co-operating with a blast, was productive of the congelation of water. Pure prussic acid will enable me hereafter to exhibit a phenomenon still more surprising; I mean that of the freezing of one portion of a liquid, by the vaporization canother portion. I shall now proceed to show that the freezing of water may be caused by the ebullition of ether

Water Frozen by Boiling Ether.

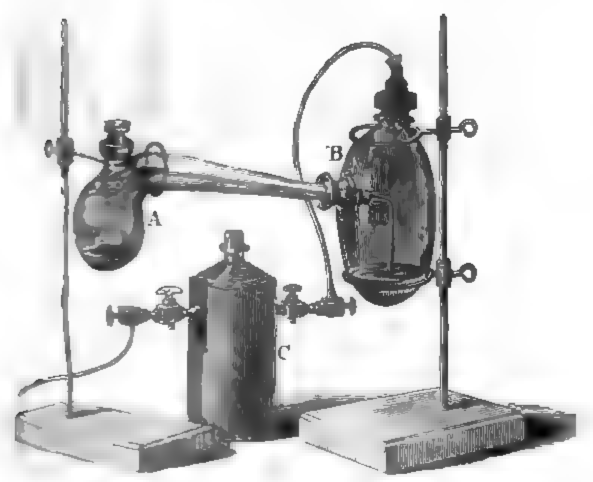
396. Let a portion of water, just adequate to cover the bottom, be it troduced into the vessel represented in the following engraving, as surpended within a receiver. Over the water let ether be added, in quartity sufficient to form a stratum from an eighth to a quarter of an inc in depth. If, under these circumstances, the receiver be placed on the air pump plate, and sufficiently exhausted, the water freezes, while the ethe boils.





397. Rationale.—The freezing of the water in contact with the boiling ether, is in consequence of that increased capacity to combine with caloric already explained. (186.) Under these circumstances, the boiling point of the other is depressed below the freezing point of water; and consequently it causes the congelation of that liquid from the same cause, that melted tin or lead will congeal under boiling water.

Engraving and Description of an Apparatus and Process for the rapid Congelation of Water, by the explosive Evolution of Ethereal Vapour consequent to the combined influence of Rarefaction, and the absorbing power of Sulphuric Acid.*



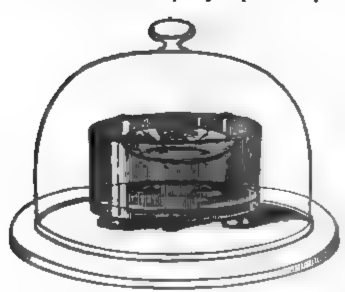
The report A, contains a small portion of water covered by a sum of hydre sulphuric other. The vessel B, holds a stratum of sulby acid of about two nucles deep, at the deepest part. Into a tubulure the ade of this vessel, the beak of the retort is ground to fit air-tight, and

^{*} The Beraldy of the American Philosophical Secrety, I am permitted to introte to article in my Compendant, although a minumented to them for a volume [her Instructions now in the press

is made to receive one end of a recurved tube, of which the other scends about half an inch below the surface of the acid. There is cury bottle, C, of which the mouth is well closed, and which is furnish two cocks, one of which communicates with the air pump, the other vessel, B. The mode of operating is as follows: the bottle is prexhausted, and kept in a state of exhaustion by closing both of the pump being put into operation and the cocks opened simultaneor power of the acid to absorb the vapour, co-operating with that of cum and the pump in exhausting the air and vapour from the retorn an explosive vaporization of the ether, and a consequent rapid cot of the water.

Congelation of Water in an exhausted Receiver by the aid of St Acid.

399. In the experiment above illustrated, water is frozen by the abstraction of caloric, consequent to the copious vaporization of eth unrestrained by atmospheric pressure. In vacuo, water undergoes rization, analogous to that of the ether in the preceding experiment; aqueous vapour evolved in this case is so rare, that it cannot act the air-pump valves with sufficient force, to allow of its being pumps a receiver with the rapidity requisite to produce congelation. How



the process which I at to describe, water r frozen by its own vapor 400. A thin dish,

of glass, covered by quantity of water, and over some concentrat phuric acid in a broat is placed within a r on the air-pump plate presented in the annergraving. Under these stances, the exhaustion receiver causes the con of the water.

401. Rationale.—So long as there is no diminution of the thin vapour which, in the absence of the air, occupies the cavity of the t the elastic reaction of that vapour prevents the production of more but when, as in the case in point, the vapour is largely in contact w phuric acid and consequently rapidly absorbed, a corresponding vition of the water takes place to supply the deficiency thus created caloric requisite for the generation of the vapour thus formed, is take the residual liquid, which finally freezes in consequence. (229.)

Improved Apparatus for freezing Water by the aid of Sulphuric

402. Finding the experiment, for which the apparatus represente preceding figure is usually employed, liable to fail from the imperfecces, dependent for their efficacy on a metallic joint, I contrived the ratus which the opposite engraving is intended to represent, and shall proceed to describe. A brass cover is so well fitted to the r

Apparatus for the Congelation of Water in Vacuo, by means of Sulphuric Acid.



(Page 70.)

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sered with supraise accu, and another jar with feet, also supplied at enough to make a stratum half an inch deep on the bottom, is instant feet, kept at such a height above the surface of the acid in the ar, as not to touch it. Upon the surface of the glass vessel, a small very thin sheet brass is placed, made concave in the middle, so as a small quantity of water. The brass cover is furnished with three pocks, one communicating with the air-pump, another with a barometge, and the third with a funnel supplied with water.

With the apparatus thus arranged, having made a vacuum on a lay, I was enabled to freeze water situated on the plate, and to keep a congelation till the Thursday following. As water in the state of reporates probably as fast as when liquid, the whole quantity frozen I have entirely disappeared during the night, but for the assistance of the mass of ice was at times about two inches square, and from ter to a half an inch thick. The gradual introduction of the water, i of the funnel and valve cock, and of the pipe represented in the by which it was conducted to the cavity in the sheet brass, enabled accumulate a much larger mass than I could have otherwise prolate the straps proceeding from it, serves to keep this jar in a proper positiat is, concentric with the outer jar.

4. In this experiment, I employed an air-pump upon a new construcwhich I contrived a few years ago, and of which a description will be

a in the Appendix.

Congelation, as effected in the experiments above described, may be maked by the aid of any substance having a very strong affinity for instance chloride of calcium, clay, or whinstone, after having andered anhydrous by ignition. Even parched meal or flour has accessfully employed in the process.

Of the Freezing of Mercury by the Vaporization of Ice.

196. If a pear-shaped mass of ice containing the metal be suspended in a large surface of sulphuric acid, and a good exhaustion obtained, it freezethe quicksilver which may be kept solid for several hours.

Wollaston's Cryophorus.

407. The adjoining figure represents the cryophorus, or frost bearer, an instrument invented by the celebrated Wollaston, in which congelation is produced in one cavity by rapid condensation in another, consequent to refrigeration.

409. In form, this instrument obviously differs but little from the palm glass, already described. (213,&c.) It is supplied by the same process with a small portion of water instead of alcohol; so that there is nothing included in it but water,

either liquid or in vapour.

409. The cryophorus being thus made, if all the water be allowed to run into the bulb near the bent part of the tube, and the other bulb be immersed in a freezing mixture, the water will be frozen in a few minutes.

410. Rationale.—There is no difference between the causes of thi nomenon and those by which the congelation of water in vacuo is e by the aid of sulphuric acid; excepting that in the one case the ac vapour is absorbed by the acid, in the other condensed by the cole either instance it is rapidly removed, and a proportionably rapid vartion of the water ensues, abstracting the caloric of fluidity from the reportion.

Large Cryophorus.



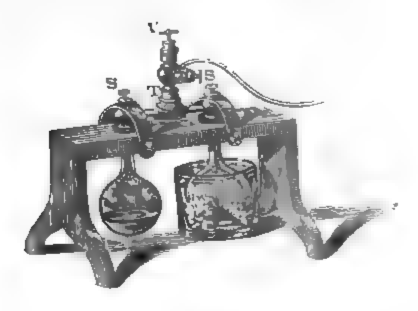
411. This figure represents a very large cryophorus, the blowi which I superintended, and by means of which I have successfully rep Wollaston's experiment.

412. This instrument was about four feet long, with bulbs of about

inches in diameter.

Modification of the Cryophorus.

418. Two flasks, of which the necks have flanged orifices, are so so in a wooden frame that, by the pressure of screws, S S, and gum of disks, the orifices of a tube are made to form with them severally, an junctures. The orifices of the tube are furnished with brass flanges, correspond with those terminating the necks of the flasks.



414. Midway between the junctures a female screw is soldered tube for the insertion of a valve cock V, by means of which, and a fl tube extending to an air-pump, the flasks may be exhausted, and closed. A small quantity of water having been previously introduce one of them, if, while the exhaustion is sustained, the other flask be rerated by ice and salt, the water will be frozen.

415. This apparatus may be applied to the purpose of desiccatiplacing the article to be dried in one receptacle, and quicklime, chlocalcium, or concentrated sulphuric acid in the other. The orifice receptacles may be made larger without inconvenience. Two large

ders, for instance, may be used.

Chemical Combination as a Cause of Cold.

M. Chemical union, although more frequently the cause of increased puttre, is in many cases productive of the opposite effect.

If There are few instances of chemical union, which are not accomilly a change of capacity. Of the cause of such changes, we are hignorant, and of course have no more reason to wonder when, by an pion of caloric, cold is the consequence of chemical reaction, than , by an evolution of caloric, heat arises from the same source.

is in the case of the solution of snow in concentrated sulphuric acid, by address, we find these opposite effects resulting apparently from the cause. Under the same head of solution, as a cause of heat or cold, it remined that nitre and nitrate of ammonia produce cold during their sometimes of artificial cold, is the solution of ice, in consequence of the

m between it and the more deliquescent salts, or the mineral acids. I It may be inferred, from the statements already made, that the mure of freezing water, or melting ice, is 32°; and that when ice is nded by other bodies at a higher temperature, it will continue to abfrom them the caloric necessary to its fusion, until it be all liquefied. t be evident that the minimum temperature which can be thus attain-**But** by mingling ice in a divided state, with certain salts or having a great affinity for water, and which form with it compounds the feezing point is lower than 32°, the mass will abstract caloric fining bodies in a mode quite analogous to that in which ice has med to operate; while the minimum temperature attainable is as wer as the feezing point is lower. Thus the freezing point of salt is about zero of Fahrenheit's scale; consequently on mingling snow, the fiquefaction of the resulting mass will proceed, at any mare above zero, to abstract caloric from all adjoining bodies until sas cold as the mixture. By the addition of crystallized chloride zm. or of diluted nitric or sulphuric acid, to snow, a compound may med, of which the freezing point is below that at which mercury . or-39°. Housekeepers have latterly availed themselves of the e of salt, to remove ice from the marble steps at the entrance of wings; as in this way it may it may be detached without injury to

Table of Freezing Mixtures.

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The following tables are taken from Thomson's Outline of the sof Heat and Electricity, page 191.

Frigorific Mixtures without Ice.							
Mulgre.				Parts.	Thermometer sinks,	Degree of a cold pro- duced.)
f amenia	•	•	•	- 1 }	From $+50^{\circ}$ to $+4^{\circ}$.	46	ì
famonia b of mda	•	•	•	- }	From $+50^{\circ}$ to -7° .	57	ر ب
b of mda pittic scid	•	•	•	- 9 \ - 4 \	From $+50^{\circ}$ to -12° .	62	7
Connecia Connecia Delt seid	•	•	•	- 9 - 6 - 4	From $+50^{\circ}$ to -21° .	71	
i min	•	•	•	- 8} - 5}	From $+50^{\circ}$ to 0° .	50	}
saffaric aci	ď.	•	•	- 5 } - 4 }	From $+50^{\circ}$ to $+3^{\circ}$.	47	<i>!</i>

Frigorific Mixtures with Ice.

Mixtures.				Thermometer sinks, Parts.	Di co di
Snow, or pounded ice Muriate of soda		•	•	$\left\{\begin{array}{c} 2\\1\end{array}\right\}$ From any temp. to -5° .	
Snow, or pounded ice	•	-	•	- 12)	
Muriate of soda	•	•	•	- 5 From any temp. to — 25°.	
Nitrate of ammonia Snow	•	•	-	$\{\begin{array}{c} 5 \\ 3 \\ 2 \\ 3 \\ 3 \\ 3 \\ 3 \\ 3 \\ 3 \\ 3 \\ 3$	
Diluted sulphuric acid		•	•	• &)	
Snow Muriatic acid -	•	•	•	$\frac{8}{5}$ From $+32^{\circ}$ to -27° .	
Snow	•	•	•	$\{ \begin{array}{c} 7 \\ 4 \end{array} \}$ From $+32^{\circ}$ to -30° .	
Diluted nitric acid Snow	•	•	•	4	
Muriate of lime -	•	•	•	From $+ 32^{\circ}$ to $- 40^{\circ}$.	
Snow Cryst, muriate of lime	-	•	-	$\left\{\begin{array}{c} 2\\ 3\\ \end{array}\right\}$ From $+32^{\circ}$ to -50° .	
Snow	•	•	-	- 1 (From 00 to 660	
Cryst. muriate of lime Snow	•	•	•	· 2	
Cryst. muriate of lime		-	-	-3 rrom -40° to 73°.	
Snow Diluted sulphuric acid	l	-	-	$\begin{array}{c} . \\ . \\ . \\ . \\ . \\ . \\ . \\ . \\ . \\ . $	
•				-	

STATES IN WHICH CALORIC EXISTS IN NATURE.

- 421. With two of the modes in which caloric exists in nature, the dent of this Compendium has been made acquainted; and these are the modes of its existence generally recognised. As it exists in one of the is called sensible heat, being susceptible of detection by the senses, or the thermometer. In the other it is called latent heat, because the quantity sent in that mode of existence, is not open to those means of detection. even in this latent state, caloric is known to be influenced by temperabeing liable to be removed entirely from vapours, or liquids, by come cation with colder substances; so as to render its subsequent present these, a proof of its previous existence in the matter from which it have been abstracted.
- 422. It seems to me, however, that, in some substances, caloric evidexists in a state in which it is wholly independent of external chang temperature. In this predicament I suppose it to reside in the nit chlorates, and fulminates, and generally in all detonating compounds.
- 423. If, agreeably to the received chemical doctrines, we are to a the explosive power of such compounds to combined caloric, it must be dent that its condensation in them is wonderfully great. Yet no good rean be assigned for this prodigious condensation. It cannot be assimply to the attraction of ponderable matter; since the same pondematter which confines it at one moment, liberates it in the next without adequate assignable cause.
- 424. Thus the presence of platinum sponge, a cold metallic cong causes the caloric of a gaseous mixture of hydrogen and oxygen to explosively. An electric spark, or the contact of any ignited matter duces the same result. The case of gunpowder, exploded by the ig of the most minute portion of the mass, is equally unaccountable, an wise the explosive recomposition of water by a discharge from the galvanic wires, by which its decomposition may have been effected.
 - 425. The almost irresistible extrication of oxygen in the gaseous

LIGHT. 75

hom oxygenat , by co x with the oxide of silver, is still more in

hing and even more surprising

in a peculiar state, dependent nidden cause, of which the detection would probably unfold 1 ny my ries in galvanism and electromystism, as well as in cher stry. I deem it more than probable that the cause of electricity is the principal agent in these mysterious phenomena.

SECTION II.

LIGHT.

427. It must necessarily belong to chemistry to treat of light, so far as it is productive of heat, deoxydizement, and other chemical effects, and so far as it is evolved by chemical processes.

438. According to Ne rton, light is a subtile fluid, which is either radiated or ref cted from every visible point in the miverse, in conseque of its elasticity or the self-

mellant power of its pa tie

429. It comes from the about ninety-five millions of miles, in eight minut 1 ly at the rate of two handed thousand miles in a seco

- 436. Light appears to have no sensible weight. The products of the combustion of phosphorus, carbon, and other combustibles, app ar fully equal in weight to the productable matter employed. It follows that the loss of the light and heat occasions no diminution of weight; yet enough is emitted by the flame of a candle or lamp to be perceived by many hundred millions of eyes. There is not a luminous point in the universe, from which a sphere of says is not emitted, in radius equal to any distance from which that point may be seen.
- 431. According to Huygens, Euler, Young. Fresud, and others, light is the undulations of a rare elastic medium, or ether, which pervades to universe. This opinion has, within the last forty years, gained the application of a majority of men of science. The doctrine of Newton is, the difficult to comprehend, and serves sufficiently to associate the particular of calorific repulsion, (11, &c.) we cannot consistently explain the communication of heat (289, &c.) without supposing that the particular radiate from hot bodies, as do those of light from luminous that agrees bly to the Newtonian doctrine. But if calorific radiation be instabled the emission of material particles by hot bodies, it would be in-

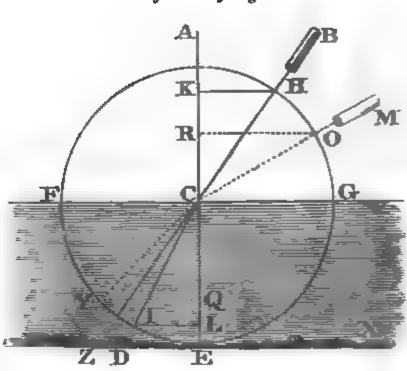
consistent not to ascribe the analogous phenomena of light to a like c In obedience to these considerations I shall resort to this theory in tro of light as a chemical agent, not without a hope that the objections thave been made to it, may hereafter find an answer in some new view c subject.

Of the Sources of Light.

432. As a source of light, the sun is obviously emore prolific than as a source of heat; and it mus evident, that all the processes which produce ignition also produce light.

433. There are some cases in which light is emi without heat. As it comes to us from the moon, as e ted by luminous insects, decayed wood, or the phospho cent wave, it appears to be unaccompanied by caloric.

434. In the fire-fly, and in many other insects, i evolved by vital action.



Refraction of Light.

435. When a ray of light passes obliquely from a rarer into a denser medium bent towards the perpendicular direction. When the course of the oblique from the denser medium into one which is rarer, it is bent from the perpend direction.

436. Suppose F G X Z to be a body of water. If a pencil of the solar ray upon the surface of the water perpendicularly at C, it will penetrate the water out deviating from its previous course, for whatever may be the attraction be the light and the water, it cannot cause any deflection, since it must act equa either side of each ray. But should a pencil of rays passing through the tu and penetrating the water at C, reach the bottom, it would shine on the pebh whereas, it would shine upon Z, were the water removed. The light in thi passing from a rarer into a denser medium, and entering the latter obliquely, there attracted by the denser medium most on the side nearest to it, and consequence bent, or refracted, from their previous course.

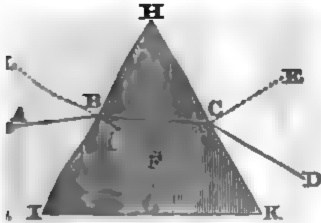
LIGHT. 77

Most C, as a centre, describe the circle, F H E, and from A draw a diameter, perpendicular to the surface of the water. Let the lines B C, C I, represents of the light in passing from the tube to the bottom of the water. these lines intersect the circle, draw K H, I L, parallel to the surface of the The angle A C H, which the incident ray makes with the perpendicular, I the angle of incidence, and K H the sine of this angle. I C E is called to differentian, and I L its sine. In the case of water, the sine I L is alread to be the sine K H, as 3 to 4; but were a mase of glass substituted for m, the sine of the angle of refraction to that of incidence would be as 2 to 3, he glass were replaced by a similar mass of diamond, the ratio would be a 2 to 5: the ratio being always invariable in the same medium, whatever in of incidence may be; for if the pencil of rays were to proceed to C, from at M, making the angle of incidence, A C M, and the angle of refraction, the size, Y Q, would be the sine, R O, in the same ratio as I L to K H; and ald half good as before stated, whether F G X Z were water, diamond, cryster born specially ensures then light passes out of such madia into the wis it is air alone that it takes place; it is enough that the substances through a passes be of different densities, or chemically different in their natures. Able Equids or colids have been found to refract most powerfully. It was his my of the association between combustibility and refracting power, that led at Newton truly to infer the combustible nature of the diamond, from its sufficient in causing refraction.

As an illustration of the case of light refracted, in passing out of denser into sour, let us imagine the eye of an observer placed at the upper orifice of to, B, in the figure. Instead of the pubble, Z, which he would see if the water moved, the pubble, D, will be seen by him. Hence the well known power to in readering an object visible, when, in the absence of the liquid, our view to intercepted by the side of the containing vessel; and hence likewise the simage which a nick or cord presents to us, when seen partially under water.

was between the Befracting Influence of a Triangular Prism, and of a Plate or Pane of Glass.

In passing through a plate of glass whose surfaces are parallel, the refraction a light sustains from one surface, is compensated by an opposite refraction by the surface; but during its passage through a prism as represented in the following ram, it is subjected to a concurrent refraction from two surfaces.



440. Supposing that the refracting medium, F G X Z, in the last figure, were bounded by air below as well as above, and its upper and lower surfaces were parallel, as in the case of a plate or pane of glass, a ray of light in passing obliquely through it, would be equally attracted, on one side, as it emerges. Hence, after its emergence, it will proceed parallel to its original direction.

41. But should a ray fall uption, as represented in the foregoing figure, in the direction of the line, A B; My to the preceding demonstration, it will, on account of the obliquity of its th, to refracted towards C, and emerging from C, obliquely to another surtice prisms, B C K, it will be again most attracted by that portion of the surments which it inclines Consequently, it will be refracted so as to proceed firection C D.

Then it must be evident that the two surfaces of the prism have a concurrent in handing the rays from their previous course; while in the pane, the in-

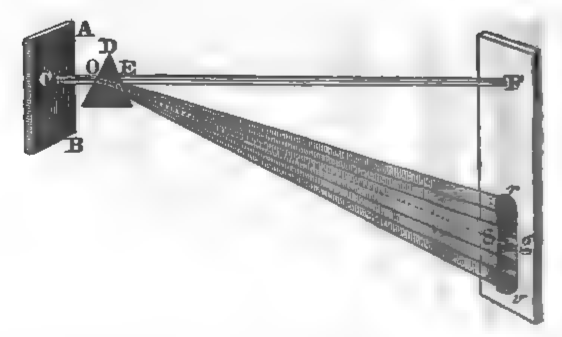
the lines, L F and E F, being perpendiculars to the surfaces of the prism, the single of incidence, and F B C, the angle of refraction, to the surface the says enter the prism. F C B is the angle of incidence, and E C D, the incidence to the surface from which the rays emerge

Dispersion of Light.

444. Besides the refraction sustained by a pencil of rays, agreeably to the pen t

ceding illustration, they undergo another alteration, the effects of which are very pleasing, and, agreeably to the doctrine of Newton, highly instructive, being the foundation of his theory of colours.

445. Light appears to consist of particles of different kinds; each kind having the property of producing on the retina of the eye a peculiar impression, which heir conveyed to the sensorium creates the idea of a colour. The rays thus capable of acting differently on the retina grown to be appeared to the sensorium creates. of acting differently on the retina, seem to be unequally susceptible of refraction. Hence, in passing through the prism, they are separated from each other, forming abeautiful series of all the various colours of the rambow, in an oblong figure called the spectrum. Under these circumstances, the rays are said to be dispersed, and the process by which they are separated is called dispersion.



446. Let A B represent a part of a window shutter of a room, into which high enters only through the hole C. If the light thus entering be received on a screen, a circular spot on it will be made luminous. But if a glass prism, D O E, be placed before the hole, so that the light may fall advantageously upon the prism, the rays, which had before produced the luminous circle, will be refracted and daspersed. as to form the spectrum, r g v, consisting of the following colours, arranged in the following order—red, orange, yellow, green, blue, indigo, violet.

Of the Heating, Illuminating, and Chemical Properties of the Rays.

447. The red rays are found to be pre-eminent in heating power; the violet as remarkable for their superior influence in certain chemical changes, dependent on deoxidation. In the middle of the spectrum, the rays have the highest power of illumination.

448. Besides the rays thus mentioned, there are invisible, heat-producing rays beyond the red, and invisible rays

producing deoxidation beyond the violet-

449. Agreeably to the observations of Herschel, to whom we are indebted for the discovery of these invisible rays, the greatest heating and deoxidizing power exists just beyond the limits of the visible spectrum; but from

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by Seebeck and Mellone, it appears that the beating of the principal heating power is dependent

m in mature of the refracting medium.

450. In the spectrum pro he principal heat was in t peta having been made blow glass prisms, it v eccupied by water or ale te yellow rays; when it solutions of sal-ammonia un heat was in the o

ed by crown or plate glass, red, and in that procured by firt glass, beyond the red; variety of transparent liquid y the cavities of several that when a prism was the maximum of heat was in is filled with sulphuric acid, or or corrosive sublimate, the max-

451. Of the rays per x by the eye, the red, being in the the least bent from the r vious course, are obviously the least refrangible; a it is no less obvious that the violet, being the most b it, are the most refrangible; also that those rays, which are found equidistant from the red,

mi violet, have a mean refrangibility.

452. An opinion has en entertained by some philosoher that there are only three original and distinct species dight, which seems lately to be sanctioned by one of the mest celebrated opticians of modern times. I allude to Sir David Brewster, whose opinions I shall give, by quoting them in his own language, from his Treatise upon Optics, page 68, American edition.

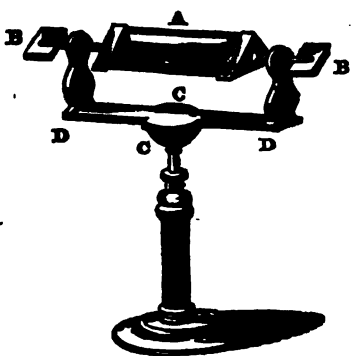
With the view of obtaining a complete analysis of the spectrum, I have connect the spectra produced by various bodies, and the changes which they unby absorption when viewed through various coloured media, and I find that the colour of every part of the spectrum may be changed not only in intensity, but modow, by the action of particular media; and from these observations, which it well be eat of place here to detail, I conclude that the solar spectrum consists of three spectra of equal lengths, viz. a red spectrum, a yellow spectrum, and a blue spectrum. The primary red spectrum has its maximum of intensity about the midde of the nd space in the solar spectrum, the primary yellow spectrum has its maxmem in the middle of the yellow space, and the primary blue spectrum has its max. between the Was and the indigo space. The two minima of each of the three process spectra, coincide at the two extremities of the solar spectrum.

- From this view of the constitution of the solar spectrum we may draw the Moving conclusions:-

-1. Red. yellow, and blue light exist at every point of the solar spectrum. 2. As a certain portion of red, yellow, and blue constitute white light, the codevery point of the spectrum may be considered as consisting of the predomi-Thing colour at any point mixed with white light. In the red space there is more when is necessary to make white light with the small portions of yellow and blue which exact there; in the yellow space there is more yellow than is necessary to white light with the red and blue; and in the part of the blue space which ap-Provided there is more red than yellow, and hence the excess of red forms a violet to blue.

E-3. By absorbing the excess of any colour at any point of the spectrum above But the same ry to form white light, we may actually cause white light to appear that the and this white light will possess the remarkable property of remaining the any number of refractions, and of being decomposable only by absorption. Such a white light I have succeeded in developing in different parts of t These views harmonize in a remarkable manner with the hypothesis of t which has been adopted by many philosophers, and which others had a its incompatibility with the phenomena of the spectrum."

Triangular Glass Prism, conveniently mounted on a univers



This figure represents a glass prism, mounted on B joint, supported by a bras as to be well qualified for sion of light, agreeably to ments alluded to in the pr ticles.

A, the glass prism, st each end by a pivot.

B, B, handles by mear the pivots are turned, so the prism revolve.

C, C, ball and socket, joint, upon which the plate

be moved so as to assume any serviceable position.

Of certain Chemical Effects of Light.

458. I have already adverted to the calorific in light, and to its power of producing chemical Among these, the bleaching power of the solar I miliar to every body. In this process the rays a exercise that modifying influence on the attractio derable matter already alluded to. (20, 21.) Con a new arrangement of particles ensues in lieu of the formed the colouring matter. Certain vegetable exposed to the sun in water, have been found to y gen gas. Some metallic salts, especially nitrate are blackened by exposure to light, owing, as is a deoxydizement. A mixture of hydrogen and chlori the dark, remain for a long time without combi in the rays of the sun will explode. According lius, the power of producing this result exists o violet rays.

459. Other important processes in which chem tion is produced by the agency of light, will be a as I proceed.

Polarization of Light.

460. This name has been given to a property which causes it often to be divided into two por

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which is translated, the other reflected, by the same glass: or one portion sustains refraction in an orange degree, the other in an extraordinary degree. Again, here properties are found to be commutable, so that portion of the rays which is reflected in one case, may be the ordinary refraction, in another may undergo the the ordinary refraction, and vice versa.

These phenomena are ascribed to the different mitical animal by the different groups of rays, in conceptate of which certain poles, which the lumeniferous mittees as supposed to possess, are variously directed at times, so as to determine their reflection, or trans-

or the degree of their refraction.

In consequence of this diversity of position, in the public light-producing particles, and the peculiar arrangement of the particles of certain transparent bodies, those lights of light, of which the poles are favourably situated transmission, may pass through such bodies, when the portions, of which the polar positions are different, we reflected; one group of the rays may undergo transmission, reflection, or refraction, the polarity of the goups of rays being reversed, those which were manitted, or unusually refracted, in the first instance, my, in the second, be reflected at first, or ordinarily facted, may, in the second, pass through, or be unusually facted.

these of Edinburgh, that the non-luminous rays emitted heated bodies, are susceptible of affections analogous these ascribed to the polarization of light. As the manera in question are due to the reaction which takes between masses and particles, agreeably to the deficion at the commencement of this work, they belong to mal philosophy proper, not to chemistry. Yet a chemist heat be indifferent to inquiries which tend to sanction, or that, his theoretic deductions respecting the important minerating phenomena of heat.

OF PONDERABLE MATTER.

464. Whatever may be the real state of the case, it has been found con venient by chemists, during the last forty years, to assume the existence of three imponderable principles, in order to account for certain phenomena and associate them advantageously. The reasoning which tends to justify this course, has been already briefly stated. (10 to 22.) Of two of those principles, calcric and light, I have treated in the preceding pages. Of the other imponderable principle, electricity, whether statical or dynamic separate treatises will be supplied.

465. In the next place, I shall treat of that "kind of matter which is endowed with weight, and which is in consequence recognised as materia by the mass of mankind." (18.) This kind of matter may be generically

designated as ponderable.

designate substances, which are exclusively or generally the products of animal and vegetable organization, as organic, all other matter being designated as inorganic. Hence, nominally, two branches of chemistry have been created, called organic, or inorganic, accordingly as the objects of electron have been such as to justify the one, or the other designation. Yet it is undeniable that no accurate line of demarcation can be drawn between the branches thus distinguished. Substances produced by animal or vegetable life, may in several instances be obtained by the reaction of inorganic bodies; the phenomena in each branch are dependent on the same ultimate elements; and in almost all cases, those of organic chemistry are displayed by means of agents derived from the inorganic world.

467. Nevertheless, the separation of chemical science into the two branches in question, seems to me highly advantageous in practice. Few persons who are not chemists by profession, can acquire more than a general chemical knowledge of important facts, properties, elements, principles, and combinations, with so much theory as may be necessary to associate them. With those details and minutiae, of which organic chemistry mostly consists it were useless to endeavour to impart a knowledge during the time allotted to an education, in which the attention of the learner is divided between seve ral branches of science. But the acquisition of that degree of knowledge which it is reasonable to expect in organic chemistry, is quite easy to a student who is familiar with the inorganic department of this science; while to one ignorant of the latter, the smallest progress in the former is utterly

impracticable.

46%. This subject will be recurred to when I enter upon organic chemis try. Meanwhile, after treating of certain general properties of ponderable matter, or the means of ascertaining or observing them, I shall process with the chemistry of inorganic substances.

OF CERTAIN GENERAL PROPERTIES OF PONDERABLE MATTER,

AND OF THE MEANS OF ASCERTAINING, OR OBSERVING THEM.

469. As introductory to the consideration of the individual inorganic substances, it will be expedient to treat of Chemical Attraction, Definite Proportions, Specific Gravity, and the Mode of collecting and preserving Gases, formerly resignated as Pneumatic Chemistry. These subjects will be considered in the four following sections.

SECTION I.

OF CHEMICAL ATTRACTION.

ttraction which takes place between heterogeneous partides only. I object to this restriction of its meaning, betance I consider it as affording a natural line of separation between chemical and mechanical philosophy, to consider the one as treating of the reaction of masses, or of masses and particles, the other of the reaction of particles only. Besides, the process of crystallization, of which I shall in the text place treat, arises from the reaction of homogeneous atoms; and it was among chemists that the investigation or observation of the laws and phenomena of crystallization originated. I consider the force which causes homogeneous atoms to cohere, whether in the crystalline form or otherwise, as a species of chemical attraction.

471. The attraction which takes place between homogeneous particles, is designated as attraction of aggregation, extraction of cohesion, or homogeneous attraction. The attraction which arises between heterogeneous particles, is

called chemical affinity, or heterogeneous attraction.

Of Attraction of Aggregation or Cohesion, or Homogeneous
Attraction. -

472. Of this kind is the force which enables bodies to re-

the tae word particle only to designate those elementary portions of matter which want by any natural means be divided. Chemists use the word atom to their such a particle, or the smallest portion of a chemical compound, which cas exist without decomposition. (472, 507, 550, 551.)

chemical nature of a substance. It is the cause of crys lization. (See note.)

Of Crystallization.*

473. Almost all matter, in passing from the fluid to solid state, assumes regular forms called crystals. inconceivable that homogeneous particles, or atoms, differ in size or shape, it is not wonderful that wl united by the same attractive force, they should produ regular forms. To produce irregular forms, the atoms, the forces actuating them, should be irregular. In fact, the deposition of matter from solution, or on the evapo tion of the solvent, is accelerated or retarded, a corn ponding change ensues in the crystalline form. way various deviations arise from that primary form wh is assumed under circumstances which allow the deposit to proceed at the same rate precisely. Those forms, wh deviate from the primary form, are called secondary. I various steps by which they are generated from the mary forms, have been most ingeniously traced, or inferr by Hauy and others. In some instances, the primary primitive form has been developed by cleavage.

474. It was at one time the general impression, t every chemical compound had an appropriate crystall form. Latterly it has been shown that certain substan quite different in their nature, as for instance, phosphc and arsenic acid, assume the same forms in crystallizi Such substances are said to be isomorphous. In the int duction to Thomson's Inorganic Chemistry, several grown

of isomorphous substances are mentioned.

475. Other things being equal, crystals are larger proportion as their growth is slower. They shoot frextraneous bodies, as the sides of the receptacle, or frestrings or sticks, in preference to crystallizing in an isoted manner. Agitation hastens their production but cofuses them. The crystalline texture of some of the trocks is attributed to slow cooling. The same matfused, and allowed less time to cool, forms a glass.

^{*} The details of crystallography, as they have been presented by Hauy and ot are of themselves so copious as to require for their remembrance a greater effethe mind than all the chemistry which I expect a candidate for a medical degracquire. It is evidently one of those subjects of which a copious knowledge cabe imparted advantageously during a strictly medical education. The instrument of the instrument of the instrument of the extremely brief.

that, if two flasks, both containing mated solution of two parts of nitrate of potash and a sulphate of soda, be surrounded with ice or cold, on introducing a crystal of nitrate of potash into and a crystal of sulphate of soda into the other, als will be formed in each flask, of the same nature at of the crystal introduced. Nitrate of potash will und crystallized exclusively in the flask first mend, and sulphate of soda as exclusively in the other.

7. Crystals are found in nature and are produced ar-

dy.

8. The precious stones e native crystals. Carboof line, common salt, and gypsum, are native prooften crystalline in form.

1 Gonismeter, or Angle Measure; on Instrument for measuring the Angles of Crystals.

Crystals may appear to be exactly similar to the eye; but when compared by accurate instruments called goniometers, they will often be found to differ rangles. Of these instruments there are two constructions; one, being more the weed, is of more general utility; the other, contrived by Wollaston, is used, but when skilfully employed is capable of giving more accurate re-

The instrument of the ensiest application, and which is usually employed, is usually the following engraving.

B B B

Of the Common Gonismeter.

construction is founded upon the 15th proposition of Euclid, which defect the opposite angles, made by any two lines in crossing each other, lines it fellows that the angles made by the legs, B B, B C B, of this choice and below the pivot on which they revolve, are equal to each y be made to close upon any solid crystalline angle, president at C, the contribution of Euclid, which describes any two lines in crossing each other, and the contribution of Euclid, which describes the contribution of Euclidean Allow the provide any two lines in crossing each other, and the contribution of Euclidean Allow the provide any two lines in crossing each other, and the contribution of Euclidean Allow the contribution of Euclidean Allow the contribution of Euclidean Allow the provide any two lines in crossing each other, and the contribution of Euclidean Allow the Contribution of Euclidean

and is ascertained by inspecting the semicircle, A, graduated into 180 degrees, pe

cisely in the same manner as a protractor.

482. The construction of goniometers is usually such as to allow the legs to I detached from the arch, in order to facilitate their application to crystalline angle and yet, so that they may be reapplied to the semicircle, without deranging the from the angle to which they may have been adjusted.

483. The piece of brass, in which the pivot is fastened, slides in a slit in each leg so as to permit them to be made of a suitable length, on the side on which the cry

tal is applied.

Of Wollaston's Goniometer.

484. The process by which angles are ascertained by means of Wollaston's goni

meter is as follows:---

485. The crystal to be examined is attached to an axis, and so adjusted, by mear of suitable mechanism, that the image of a window bar may be seen reflected from one of the crystalline faces, so as to coincide with a line (seen directly) drawn of the wall under the window, parallel to the window bar. By a partial revolution of the axis, and consequently of the crystal, a similar coincidence of the images of the bar and line is produced by means of another face of the crystal, being the next that first employed.

486. Meanwhile the number of degrees of a circle moved through, in changing the crystal from the first to the second position, is measured by an index on a graduated arch, and the degrees of the angle, which the surfaces make with each other.

thus ascertained.

Various Modes of causing Artificial Crystallization.

487. Fusion followed by congelation.—Instances: Crystal lized sulphur, bismuth, antimony, zinc.

488. Solution followed by evaporation in open vessels.

Exemplified by salts, acids, alkalies, sugar.

489. Solution with heat followed by refrigeration.—Most of the substances which crystallize by evaporation, yield crystals in this way.

490. Solution followed by vaporization at the boiling heat.—Crystals may be thus obtained from many salts, but are

always minute.

491. Solution followed by saturation.—Instances: Potask

saturated by carbonic acid or chlorine.

492. Sublimation.—This comprises the idea of vaporization, and condensation into a state of solidity. Instances: Corrosive sublimate, calomel, iodine, arsenic.

493. Solution followed by precipitation; as in the case o

the arbor Dianæ and arbor Saturni.

Crystalline Specimens exhibited.

494. A wooden arch, about fifteen inches high and a foo wide, encrusted with fine blue crystals of the sulphate copper: also baskets constructed of bonnet-wire, curiously studded with elegant crystals of the same salt. Crystal of the ferroprussiate of potash, more properly called cy anoferite of potassium, suspended by a cord on which the

d during their formation. (475.)

There crystals are of an agreeable lemon-yellow colour.

465. A crystalline congeries of alum, about a hundred make in weight. Buskets studded with crystals of the salt.

496. Large cluster of crystallized borax.

497. Crystals of corrosive sublimate and calomel.

498. Crystals of sulphur, arsenic, bismuth, antimony, &c.

499. Various other crystalline bodies.

Of Decrystallization.

It has been ascertained by Dr. Daniell that crystals my be partially developed by solution. When alum is daily dissolved, its crystalline structure becomes very milest.

301. Specimen of decrystallized alum.

Of Water of Crystallization.

502. The well known spiculæ, which, by their appearance on the surface of water, indicate incipient freezing, are crystals. In fact, it was from the Greek name for ice, that the word crystal was adopted; as crystals were correctly considered as the products of a process analogous to freezing. This is strictly true in the case of cyttals resulting from the congelation of matter from a water enters into the constitution of many crystals which, when robbed of it by heat or desiccation, lose the crystalline form. The water thus situated E called water of crystallization. Some substances combute with water in different proportions, and consequently assume different forms; others crystallize with or without water, with a corresponding diversity of form. These reto are dependent upon variations of temperature in the polvent at the period of the crystallization. At 86°, sulphase of soda crystallizes without, at 40°, with water of cysallization. Chloride of sodium, which is ordinarily wydrous, is made to unite with water of crystallization at f below zero.

Crystals usually retain within their crevices a miportion of the solution in which they have been crysthe decrepitation of chloride of sodium
ous salts when heated, from the vapori-

zation of the water so retained. The larger the crystals, the more they are liable to this impurity.

Of the Consequence of excluding the Air from a saturated Solution of Sulphate of Soda while boiling.

504. If a flask be sealed, so as to be air-tight, while containing a boiling saturated solution of Glauber's self, (sulphate of soda,) the solution will remain liquid, so long as undisturbed, but on the admission of the air, will often become a compact crystalline mass within a few seconds. In other cases, it will continue liquid for some time, even for 24 hours, and may then crystallize on being poured out of the flask. Sometimes it crystallizes in the neck of the vessel while the operator is pouring it out; at others, allowing a crystal or other body to fall into the solution, causes crystals to shoot. No satisfactory explanation has been afforded of this phenomenon. It seems as if the repulsive and attractive powers were so nearly balanced as to enable a slight external force to determine the preponderancy in favour of the latter. That there is an evolution of caloric, consequent to the congelation, is rendered evident by a rise of temperature.

Experimental Illustration.

505. Several glass flasks being made about two-thirds full of a saturated solution of Glauber's salt, and sealed up air-tight, the solution remains liquid until the air is admitted. It then crystallizes either spontaneously, or from slight causes.

OF CHEMICAL AFFINITY, OR HETEROGENEOUS ATTRACTION.

506. This attraction is never subdued mechanically, unless when nearly balanced by repulsion; as in the case of compounds which may be exploded by percussion, (29,) or

of clastic fluids combined with liquids. (240.)

507. To sever elements, united by chemical affinity, the finest edge producible by human art is utterly incompetent. Thus, chalk consists of lime and carbonic acid; vermilion, of sulphur and mercury. Yet when reduced to powders perfectly impalpable, the minutest particle, whether of chalk or vermilion, contains the same ingredients as the mass, and in the same proportion.

Different Cases of Affinity.

50%. First Case—Simple Combination.—A and B, two exterogeneous substances, unite and form the compound A B.

Instances.

509. Copper with zinc forms brass.

510. Copper with tin forms bronze.

511 Antimony with lead forms type metal.

512. Magnesia with sulphuric acid forms Epsom salt, or suphate of magnesia.

513. Soda with sulphuric acid forms Glauber's salt, or

sulphate of soda.

514. With mercury, various metals form amalgams.

Experimental Illustration.

515. A portion of gold leaf, being triturated with mercry, disappears, forming a chemical compound with the bercary, in consequence of the inherent attraction or aftity between the heterogeneous particles.

516. Second Case of Affinity.—Called single elective at-

traction, or simple affinity.

- 517. A and B, two heterogeneous particles, being united in the compound AB, another particle, C, being blended with them in solution, unites with one of them, as A, to the exclusion of B.
- 514. In this case, C is said to decompose AB, and to have a greater affinity for A than for B.

Experimental Illustration.

- 519. Potash being added to a solution of sulphate of gresia. the magnesia precipitates in white flocks. A result takes place, on adding a solution of potash to a station of sulphate of alumina.
- 520. Retionale.—Sulphate of magnesia consists, as its implies, of sulphuric acid and magnesia. The affinity sisting between the potash and the acid being greater from combination, and, being by itself insoluble, An analogous explanation will apply in the acid alumina. In each case, the affinity of the acid

for the alkali, predominates over that of the acid for the earth.

521. Third Case of Affinity.—Called double elective at

traction, or complex affinity.

522. The compound formed by the particles A and B being blended in solution with the compound formed by C and D,—A combines with D, and B with C.

Experimental Illustration.

A B
Sulphate of zinc being mixed with Acetate of lead, C D Acetate of zinc.

A D
Sulphate of lead
Sulphate of lead
Acetate of zinc.
C B

523. Fourth Case of Affinity.—A and B being in union

C, added in excess, combines with both A and B.

524. When ammonia is added to certain solutions of metallic salts, those of copper or silver for instance, it operates at first as the potash does in the case of single elective attraction abovementioned, and the oxide of copper or silver precipitates. But if the ammonia be added in such quantity, as that, after all the acid shall have been saturated, there shall be an excess of alkali, this excess will combine with the precipitated metallic oxide, forming with it a compound which is immediately dissolved. Hence the menstruum which is at first rendered turbid, afterwards becomes clear, and, in the case of the copper, assumes to beautiful and characteristic blue colour.

Experimental Illustration.

525. Liquid ammonia being poured into a solution o copper, at first precipitates the metal in greenish flocks but, when the alkali is added in excess, these flocks disap pear, and a blue solution results.

Additional Illustrations of Chemical Affinity.

526. In order to show the wonderful power of chemical reagents in producing striking changes, some additional exemplifications of chemical affinity will here be given This exhibition may excite curiosity in the learner an

afford gratification to him, although unprepared to understand the intricate play of affinities by which the results are accomplished.

Experiments.

- 527. Silver precipitated by mercury, mercury by copper, and copper by iron.
- 528. Conversion of two liquids into an adhesive mass by mingling sulphuric acid with a solution of chloride of alcium or nitrate of lime.
- 529. Solution of ferroprussiate of potash, added to solutions of copper and iron.
- 30. Solution of chromate of potash, added to solutions of kad, mercury, and silver.
- 331. Ammoniacal nitrate of copper or silver, added to assenious acid.

Of Cohesion as an Opponent to Chemical Combination.

532. There are many substances, among others carbon, which, under certain forms, in consequence of greater hardness, are much less susceptible of chemical reaction, than under others. Thus the diamond, anthracite, charcoal and tinder, are varities of carbon, which are endowed with a susceptibility of combustion inversely as their hard-Tinder is proverbially ready to take fire, while the damond is only to be ignited by the aid of extreme heat, and as unusual supply of oxygen. Every body knows how much less susceptible of being acted upon by solvents, are bricks, porcelain, or stone ware, than the earthy matenals out of which they are made. In these cases, it would really appear that the attraction between the homogeneous stoms counteracts the heterogeneous affinity which would rer them. Yet I conceive it to be an error to confound be obstruction to chemical reaction thus created, with that which arises from the restriction of the surface in contact which solvent. Other things being equal, there will evidesity he more action in proportion as the points of conby are multiplied, and vice versa. Thus the action of will be less rapid upon a metallic ball, than upon weight of metal in the state of foil, fine wire, or traings; although the attraction of the homogeneous

particles is quite as energetic in the one case as other.

Effects of Mechanical Division experimentally illustre

533. If a ball of brass be put into one glass, and half its weight of brass filings or turnings into anoth adding nitric acid to both, a violent effervescenc ensue in the one, while in the other, the reaction will I be discernible.

Influence of Solution in promoting Chemical Reaction, rimentally illustrated.

534. Tartaric acid and a carbonate, although intin intermingled in a pulverulent state, do not react until 1 ened, when a lively effervescence ensues.

Exception to the Law that Chemical Action requires Fl experimentally illustrated.

535. If slaked lime and muriate of ammonia in p be mixed, the pungent fumes of ammonia will be perc

Tables of Affinity.

536. These consist of the names of a series of stances, placed in a column, in the order of their at for any one substance of which the name is at the he the column. The following is an example:—

Sulphuric Acid.

Baryta,
Strontia,
Potash,
Soda,
Lime,
Magnesia,
Ammonia.

SECTION II.

OF DEFINITE PROPORTIONS.

537. The proportions have been long known to be inminble, in which substances must be mixed in order to nturate each other, or to produce a compound in which the peculiar characters, or affinities of the ingredients, are stinguished.

538. When substances combine in other proportions can those of saturation, their ratio is no less definite and

constant.

539. There is not in any case, except the peculiar one of solution, an indefinite gradation in the proportions in which bodies combine. There are rarely more than four

gradutions.

- 540. The number, representing the least proportion in which a substance is known to combine, will, in a great mjority of cases, divide the numbers representing the greater proportions without a fraction; and where this result is unattainable, it will still be found that the larger proportion may be divided by the half of the lesser without a remainder.
 - Y, and Z be other substances, severally having an affinity for either A, or B, or C. Let each of the former and each of the latter be combined in the least possible proportion. Consequently, the least combining proportion of each substance will be found three times. It will appear that the proportions of A, B, and C found by combining them with X, will be in the same ratios to each other, as the proportions found by combining them with Y, or Z; and reciprocally that the proportions of X, Y, and Z, will have the same ratios, whether ascertained by their combination with A, B, or C.
 - When, instead of ascertaining the least combining proportions of six substances, the experiment has been extended to any larger number, the same uniformity has been found to prevail in the ratios of the numbers represents those proportions. It has also been found that when numbers are ascertained which express the ratio of the last combining proportions of a variety of substances to any one substance, as for instance to oxygen, those

numbers will express the ratios of the least combining portions of the substances in question, to each other

543. Numbers representing least combining propare called chemical equivalents. As they are mempressive of ratio, they may be multiplied by any comultiplier, or divided by any common divisor, v

affecting their correctness.

544. They are usually so computed as to mate equivalent of oxygen, or of hydrogen = 1. As the valents of these substances are as 1 to 8, it follows, hydrogen be represented by unity, oxygen will be oxygen be unity, hydrogen will be 0.125, or one-eigone. Consequently, equivalents, formed upon either may be converted into those corresponding with the either by multiplying or dividing by 8.

545. By Berzelius, Wollaston, and Thomson, chas been made the standard. Berzelius assumes it wollaston at 10, and Dr. Thomson at 1. The conference between the equivalents founded upon these

bers, is in the position of the decimal point.

Of Tables of Chemical Equivalents.*

546. In these, the equivalents of all known bodi far as ascertained, are arranged alphabetically. bles are of great utility in practical chemistry. rative chemist may frequently resort to them with a tage. They enable him to store his memory with adequate to the solution of a great number of que which must necessarily arise. If he wishes to know much of any two substances he must take to form a he has only to recollect, or to look for, their equivale the table, and seek a solution by the rule of three. the equivalents of the substances are to each other, the quantities of them to be used. Should it be an to produce only a certain weight of a compound, th the equivalent of the compound is to that of either ingredients, so is the weight of the compound requi the requisite weight of either ingredient.

547. In order to know how much of the proper rials he must use to effect a decomposition, he has to employ them in the ratio of their respective equiv

^{*} See Appendix for a Table of Equivalents.

48. Moreover, when the proportions, afforded by anas. b not harmonize with well ascertained equivalents, are warned of the existence of some inaccuracy, which my cases may be safely corrected so as to make the b accord with them.

Wollaston's Scale of Equivalents.

This instrument is so constructed that the computation requisite in be equivalents is performed by a slide. It has been mentioned that fivalents may be expressed in any numbers having the same ratios to ther as the least combining proportions of the substances which they the slide enables us to adopt any such numbers as may be configurable that the slide enables on the slide give the same ratios in different numbers as may be configurable to slide, we vary one equivalent to 100, for instance, requivalents vary proportionably.

Of the Atomic Theory.

Extension has been proved to be infinitely divisible, and it is not suppose that the matter, comprised within any given limits, may tible of as many subdivisions as the space in which it is contained. her hand, it is obvious, that mechanical division must be limited perfection of the edges or surfaces employed to accomplish it. Vere atoms chemically divisible ad infinitum, any one substance, mall in quantity, might be diffused, in a state of chemical combiroughout any other, having an affinity for it, however great; for perture in the latter would exercise a stronger affinity than anould be unreasonable to suppose that each should not have its hat such a diffusion is impracticable must be evident from the of the number of definite proportions to which substances in are restricted, as already mentioned when upon the subject of Hence elementary atoms are not considered as liable to an subdivision, either by chemical or mechanical agency. (539.) ratios of the equivalent numbers are supposed to be dependent entical with, those of the weights of the integrant atoms of the to which they appertain. Thus the fact that 32 parts by weight is saturate as much of any acid as 48 parts of potash, is explained log that the weights of the smallest atoms of those alkalies which are to each other on 32 to 47.

the interpose each other, no excess of either ingredient is in any rable. The line in nitrate of line is to the potash in an equivated the sulphate of potash, as 28 to 48, yet neither is the line at to take the place of the potash, nor is there too much potash to are of the line. This result is intelligible, if we suppose that, thus a just adequate for reciprocal decomposition are employed, a equal number of atoms of each salt; the one containing as not potash weighing 48, as the other contains atoms of line 38.

resame explanation applies to the fact that, while the sulphuric explanation applies to the fact that, while the sulphuric explanate of potash is to the nitric acid in the nitrate of lime as 40

to 54, yet there is neither too much of the latter: or too little of the mer, to produce neutral compounds with the bases to which they a

verally transferred.

555. On account of the hypothetical association of the numbers, senting the least proportions in which bodies are known to combine the supposed relative weight of their atoms, those numbers are a known by the appellation of atomic weights, as by that of chemical valents.

Of Chemical Symbols.

556. I shall translate from Berzelius an account of the symbols which has devised, and which it would be well to understand, as they will all met with. Objections have been made to some part of his plan, but it neral I believe it will be expedient to adhere to it; since whatever I lius recommends, awakens the attention of chemists universally, and cause his symbols to be generally understood throughout the che world.

557. "We select (says he) as symbols the initial letters of the names of bodies. When the names of several bodies have the same i we add to each a letter which it has not in common with the rest; a instance, C signifies carbon, Cl chlorine, Cr chromium, Cu copper, (balt. When, however, the names of a metallic and non-metallic elemence with the same letter, no additional letter is added to the But when two non-metallic elements have a common initial, it is need to distinguish one by means of an additional letter. Thus, to distinct chlorine, bromine, and silicon, severally, from carbon, boron, and su the symbols of the former are Cl, Br, and Si, while those of the latter

simply C, B, and S.

558. "The number of atoms is designated by cyphers. A cypher to the left multiplies all the symbols to the right, as far as the first cro or the whole formula. A little cypher, situated to the right of a sy and a little above its level, multiplies that symbol only. Thus S2O nifies one atom of hyposulphuric acid, consisting of two atoms of su and five of oxygen; while 2S2O5 signifies two atoms of the same aci such cases as that just cited, in which two atoms of the radical are with one, three, or five of oxygen, the expression for the former wo abbreviated advantageously by having a specific sign for a double The sign which I have adopted for this purpose, is a dash across the part of the symbolic letter. Thus P signifies a single atom, P2 a atom of phosphorus. Compound atoms of the first order are expressin the following example of sulphate of copper Cu O+SO3. The phate of the sesquioxide of iron would be expressed by 2Fe O3+2S659. "It may be expedient to designate the number of atoms of c

by dots placed over the letters symbolic of radicals. Thus we may nate the sulphate of copper by Cu S, the trisulphate of the sesquion

iron, by 2 Fe S3."

^{*} Instead of placing the dash across the lower part of the letter, it is go placed under it, as the former mode requires type cast for the purpose.

List of the Atomic Weights of the Simple Ponderable Substances, together with their Symbols.

As the atomic numbers are practically useful, enabling us to know produce in which substances are combined, or in which they should not produce compounds, it is advantageous to commit them to mery as far as possible. The whole number of substances recognised as matary, agreeably to the present state of our knowledge, is fifty-four. these, links more than half are of sufficiently frequent recurrence either peculation or in practice, to make it desirable to remember their number of will quote them, therefore, in two distinct tables. Those of which nowledge is likely to be rarely in demand, I have subjoined in smaller. The symbols are given in a separate column. In obedience to the symbols of potassium and sodium.

	Symb	-al			At. Wts.		QL-				At. Wts.
i ium		λų.			_	T ishin	Symbo	1.			
	Al	•	•		14	Lithium	T.	-		•	6
NOD y	Sb	•	•		64	Magnesium	Mg	-		•	12
	As	•	•		38	Mercury	Hg	-		•	202
	Ba	•	•		69	Nitrogen	N	•		•	14
th	Bi	•	•		71	Oxygen	O	•		•	8
	В	•	•		11	Phosphorus	P	_		•	16
ie .	Br	•	•		78	Platinum	Pl	•		•	99
	Ca	•	•		20	Potassium	Po	•		•	40
}	\mathbf{C}	•	•		6	Selenium	Se	•		•	40
e	M	•	•		36	Silicon	Si			•	8
	('u	•	•		32	Silver	Ag	•		•	108
•	F	•	•		18	Sodium	So	•		•	24
	Λu	-	•		200	Strontium	Sr	•		-	44
:n	H	•	•		1	Sulphur	$\tilde{\mathbf{S}}$	•		•	16
	I	•	•		126	Tellurium	\mathbf{Te}	•		•	64
	Fe	•	•		28	Tin	Sn	•		•	59
	Pb	•	•		104	Zinc	Zn	•		-	32
•	Cd			_	56	Osmium	Os			-	100
	Ce		•		46	Palladium	Pd		•	•	53
	Cr		•	-	28	Rhodium	R		-		52
	Co		•	•	30	Thorium	Th	-	-	•	60
1M	Ta		•	•	185	Titanium	Ti		•	•	24
	Ģ		-	•	18	Tungsten	W		•	•	95
_	İr		•	•	99	Uranium	Ü		•	•	217
96	Mn		-	•	24	Vanadium Vanadium	V		-	•	69
1000	Mo		-	•	4년	Yttrium	Y		-	•	32
	Ni		•	•	30	Zirconion	$oldsymbol{Z}$ r		•	-	34

appears from some experiments made by Messrs. Petit and Dulong, that the stor heat, or specific heats, of all elementary atoms are the same; so that eith heat of any one congeries of atoms be less than that of another having weight, it is because the atoms of the one being heavier than those of the reare fewer of them in the same weight. Hence the capacities, or spess, of equal volumes of elementary substances are greater, as the weights them are less; so that if, in the case of each, its atomic weight be multiplectific heat, the product will in general be so nearly the same, that the same be ascribed to the inaccuracy unavoidable in experimental investi-

hyperting this highly important and interesting inference of Petit and Duffiner A. D. Bache has endeavoured to show in an article published in the 13

Journal of the Academy of Natural Sciences, that multiplying the equivalents twolve principal metals into their specific heat, gives results so widely deviating from uniformity as to take all plausibility from the hypothesis that the atoms of simple hooling have the same specific heat

ple bodies have the same specific heat.

563. Dr. Thomson has observed that this law is more likely to be true, since holds good without doubt in the case of the gases; and that if it be true we had only to divide the specific heat of hydrogen by the atomic weight of any body, find its specific heat. Moreover that the specific heats thus found agree very near with those ascertained experimentally.

564. From the researches of Faraday, it appears that the quantity of the volts fluid given out during the solution of various metals, is in the ratio of their atom weights. It would seem, therefore, as if the imponderable atmospheres, both caloric and electricity, are held by atoms in the same equivalent proportion.

SECTION III.

OF SPECIFIC GRAVITY.

and weight are synonymous words; but the term specific gravity is use to signify the ratio of weight to bulk. Hence the object of all the processe for ascertaining specific gravities, is either to ascertain the weight of a known bulk, or the bulk of a known weight; for whether the substances whose specific gravities are to be found be reduced to the same weight and the measured, or be reduced to the same bulk and then weighed, their weights to their bulks will be discovered. If reduced to the same bulk and weighed, their specific gravities will be directly as the weights. reduced to the same weight and measured, their specific gravities will be inversely as their bulks thus ascertained.

566. Supposing a like bulk of each kind of matter in nature to be weighted, the results, numerically stated, would represent their specific gravities. But since it is not possible to procure an exactly similar bulk of each kind of matter, it is necessary to resort to another mode of reducing their bulks to a common measure. The method adopted in the case of solids and liquids, is to divide the weight of a given bulk of each body of which the specific gravity is to be found, by the weight of a like bulk of water. This in fact may be stated as the general rule for ascertaining specific gravities.

567. Thus on dividing the weight of any bulk of copper by the weight of a like bulk of water, the quotient is 9. This, therefore, is received at the specific gravity of copper. By a similar procedure, in the case of silver, the quotient is 10.5, in the case of mercury 13.6, in the case of gold 19.3: consequently, these numbers are considered as representing the specific gravities of those metals.

568. If the body be lighter than water, as in the case of cork which i only about one-fifth as heavy, the quotient, being less than one, is expressed by a decimal fraction. Thus the specific gravity of cork may the specific gravity

stated to be .2.

569. The gravity of water has been assumed as the standard, because this liquid may always be obtained sufficiently pure; and it is generall easy to ascertain the weight of a quantity of it, equal in bulk to any other body.

570. The weight of a quantity of water, equal to the body in bulk,

the resistance which the body encounters in sinking in water. z, if we can ascertain, in weight, what is necessary to overcome the mee which a body encounters in sinking in water, and divide by the k thus ascertained, the weight of the body, we shall have its specific by.

L. In the case of a body which will sink of itself, the resistance to its

is what it loses of its weight when weighed in water.

L. In the case of a body which will not sink of itself, the resistance sinking is equivalent to its own weight, added to the weight which be used to make it sink.

erimental Demonstration that the Resistance which a Body encounrs in sinking in any Liquid, is just equivalent to the Weight of a ortion of the Liquid equalling the Body in bulk.

71 This proportion may be experimentally demonstrated, by means of apparatus represented by the following figure.

574. The cylinder, represented as surrounded by the water of the vase, is made to fit the cavity of the cylinder suspended over it so exactly, that it enters the cylinder with difficulty, on account of the included air, which can only be made to pass by it slowly. It must, therefore, be evident, that the cavity of the hollow cylinder is just equal in bulk to the solid cylinder.

575. Both cylinders (suspended as seen in the figure) being counterpoised accurately upon a scale beam, let a vessel of water be placed in the situation of the vase. It must be evident, that the equiponderancy will be destroyed, since the solid cylinder will be buoyed up by the water. If water be now poured into the hollow cylinder, it will be found that, at the same moment when the cavity becomes full, the equiponderancy will be restored, and the solid cylinder sunk just below the surface of the water.

576. Hence it appears that the resistance which the solid cylinder encounters in sinking in the water, is overcome by the weight of a quantity of water equal to it in bulk. It must be evident, that the same

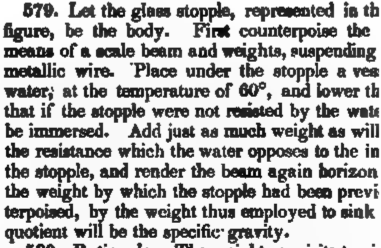
Il be true of any other body, and of any other liquid.

M7. Retionale.—When a solid body is introduced into an inelastic id, on withdrawing it a hole is left, which remains vacant of the solid have, than the liquid is found to fill up the space from which it had a removed.

within them from which they are forcibly excluded, is precisely equal weight of a quantity of the liquid commensurate with that space; when the space is reoccupied by the liquid, the equilibrium is restored. It weight, every body, introduced into a liquid, experiences from it a liquid to the weight of a quantity of the liquid, commensurate which would be produced, supposing the liquid frozen about the weight open so as to remove it, and the fragments put together the cavity thus created must obviously be exactly equal to the

bulk of the body. It follows, therefore, that the resistance whice encounters in sinking within a liquid, is equivalent to the weight tity of the liquid, equal in bulk to the body.

Method of ascertaining the Specific Gravity of a Body heav Water.



580. Rationale.—The weight requisite to suple measures the resistance to its being sunk in

and this it has been shown is equal to the weight of a bulk of wa that of the stopple. Of course, pursuant to the general rule, it cessary to see how often this weight is contained in the weight ple, to ascertain its specific gravity.

Method of ascertaining the Specific Gravity of a Body ligs.

581. Let a small glass funnel be suspended a beam, and counterpoised so as to be just below of some water in a vase, as represented in the d 582. If, while thus situated, a body lighter the small cork for instance, be thrown up under the equilibrium will be subverted. Ascertain how a will counteract the buoyancy of the cork, add weight, and divide its weight by the sum. T will be the answer.

583. Retionale.—The force with which the against the funnel, is equal to the difference weight and the weight of the bulk of water v places. Of course, ascertaining the force wi rises by using just weight enough to counter adding this weight, so ascertained, to that of t

have the weight of a bulk of water, equal to the bulk of the cor weight, dividing the weight of the cork agreeably to the gener specific gravity of the cork will be found.

Method of ascertaining the Specific Gravity of a Liq

584. Let the stopple be counterpoised, exactly as as above dimexcepting that it is unnecessary to take any account of the coweight.

585. Having, in like manner, ascertained how much weight

t given liquid, divide this by the weight required to sink it in the water.

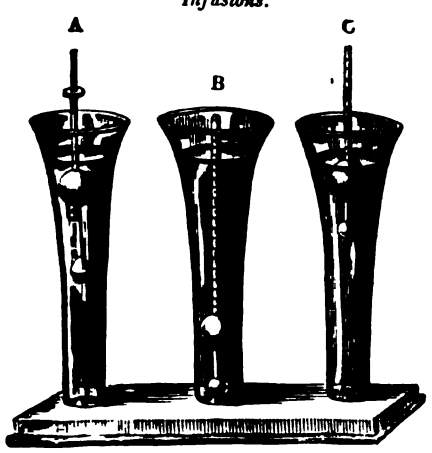
quoent will be the specific gravity sought.

Rationale.—It has been proved that the resistance to the sinking of it is any liquid, is precisely equal to the weight of a bulk of the liquid, to the bulk of the body. Ascertaining the resistance to the immersthe same body in different liquids, is, therefore, the same as ascerthe weights of bulks of those liquids, equal to the bulk of the body, if coarse, to each other. And if one of the liquids be water, diby the weight of this the weights of the others, gives their specific

If the stopple be so proportioned as to lose just one thousand grains nersion in water, division is unnecessary; as the weight of the liquid cobtained in grains, which are thousandths by the premises. A fine tal exactly of the same weight as the stopple, may be employed someonese.

In these experiments, the liquid should be as near 60° of Fahrentermometer as possible.

maters for Alcohol, for Acid, Saline, and other Solutions, and for Vegetable Infusions.



these a constant weight is used to a certain extent, and the differences of resumated by the quantity of the stem immersed. In those instruments particularly where several weights are employed, the effect is the same as med the instrument were lengthened as many times as the number of the itselfed to it.

repreceding engraving represents three hydrometers, A, B, and C, contained excis. B and C are of glass, and A of metal.

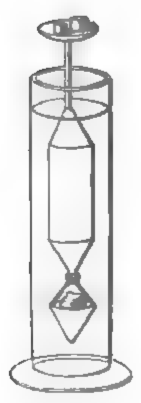
is mended for liquids heavier than water; C, for those which are lighter. he graduation commences at that point of the stem, to which the instrusivational state of the stem for liquids lighter than water, and at the bottom of the stem for liquids lighter than a the latter case, as in that of spirituous liquors or ethers, the strength ater as the lighter, more of the stem is immersed in proportion as I metronger; but the opposite is true in the case of acid and saline solutions of vegetable matter; the more the stem emerges from these, the model course the stronger they are. The instruments are represented as larger in pure water.

hydrometer of a form much used in this country and in England, both infusions of vegetable matter. The stem is virtually lengthened by Carral small weights, which may be slipped on and off at pleasure.

693. The whole difference between the weight of water and that of the spirit is equal to about two parts in ten. Of course, an hydrometer for spin have on its stem a scale of more than two hundred parts, in order to give the gravity of any liquid consisting of water and alcohol. To render such gravity of any liquid consisting of water and alcohol. To render such gravity discernible, the stem would have to be of very inconvenient. This is obviated by using different weights. When the heaviest weight is stem, the whole of the stem stands above the surface in distilled water. V liquid contains enough spirit to allow the whole of the stem to sink in it, w porting this weight, a lighter weight may be used; and when the stem against this weight, a lighter weight may be exchanged for one still Supposing the stem graduated into fifty parts, three weights would give fifty each, and the stem unloaded, fifty more. Were the stem graduated into I nineteen weights would give one hundred and ninety parts, and the stem tan more.

594. An instrument, sometimes called a saccharometer, but precisely a principle, is used for infusions of vegetable matter, especially for the wort of and distillers, excepting that the scale begins at the top of the stem, wit which coincides with the surface of pure water, at sixty degrees Fahrenhe the hydrometer is immersed in it. When the infusion is strong enough to the whole of the stem above its surface, a weight is to be added heavy esting the graduated part of the stem into the liquid. And, in like manner infusion is found stronger, weights still heavier must be added; the preceptioning analogous to, but the converse of that described in the case of size

Nicholson's Gravimeter, for accortaining the Specific Gravity of Solids, eithe or lighter than Water.



595. The accompanying cut is a representation of Rigravimeter, the construction of which is sufficiently of 596. On the upper scale of the instrument, whilst fi water, place any body, the specific gravity of which found—a piece of coin for instance—and add as much to the same scale as will sink the gravimeter, until purposely made in the stem, coincides with the surface water. The coin is then to be transferred to the low and as much weight added to the upper one as countries change. This weight is obviously just equivalence which the coin encounters in sinking in the Let this weight be called A.

597. In the next place, the body is to be removed gravimeter, and as much weight, B, again added to the scale, as will cause the mark upon the stem to saint the aqueous surface. Of the weight first employed, we need be taken; but the weight, A, and the weight B, the second and third steps of the process, are to be noted, and added together; the sum of A and B is the divided by A, the first number noted. This number presents the weight of a bulk of water, equal in the coin; while the sum of the numbers, A and B is extended to the weight of the coin; since that aggregate we been found equivalent to the weight of the coin in an gravimeter.

Method of finding the Specific Gravity of a Body lighter than Water, by M. Gravimeter.

598. Should the specific gravity of a light body, as a piece of cosk for im in question, place it on the upper scale of the gravimeter, load the install that the mark on the stem may coincide with the surface of the water, as in above stated, a leaden disk being previously laid upon the lower scale. being removed, the weight requisite to compensate its absence, gives the the cork. This weight, being added to that which will compensate its I when immersed in water by being placed beneath the leaden disk in the leading ives the weight of a quantity of water equal in bulk to the cork. How number of grains representing the weight of the cork be divided by that rep the weight of its bulk of water, the quotient will be the specific gravity, this case, must be expressed in a decimal fraction, as it is less than unity.

al of escertaining the Specific Gravity of Gascous Substances.



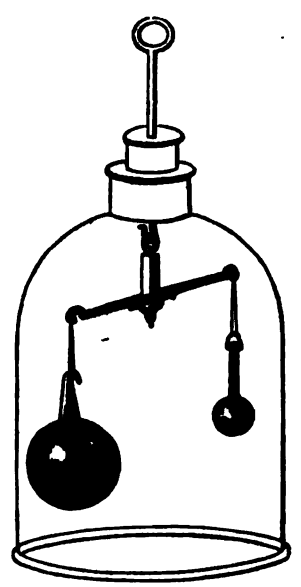
599. Suppose the globe A, represented in the adjoining figure, to be removed from the receiver, R, and exhausted during a temporary attachment to an air-pump, by means of a screw with which the globe is furnished, and which serves also to fasten it to the receiver, as represented in the figure. Being preserved in this state of exhaustion by closing the cock, let it be sus-pended from a scale beam, and accurately counterpoised, as in a former experiment. (71, &c.) In that experiment, after the giobe was counterpoised, air was admitted and caused it to preponderate decidedly. If in lieu of admitting air, the globe be restored to the situation in which it appears in this figure, so as to be filled with hydrogen from the receiver, R, and afterwards once more suspended from the beam, instead of preponderating decidedly as when air was allowed to enter, the additional weight acquired by it in consequence of the admission of the hydrogen, will scarcely be rendered perceptible. Supposing, however, that the additional weight

the same bulk of atmospheric air, after a similar exhaustion of the weights of equal volumes of hydrogen and air would be repreby the weights thus ascertained. The specific gravity of atmospheis the unit, in multiples or fractions of which the specific gravities of mare expressed. Hence the weight of a given bulk of hydrogen, by the weight of an equal bulk of air, gives the specific gravity of by a similar process, the specific gravity of any other gas may

If the Influence of the Air on the apparent Weight of Bodies.

frames illustration of the loss of weight, and consequent inaccuracy atthe ordinary process of weighing, as conducted in the air, is afforded by itself process described in the next page. (COI, &c.)

A Pound of Feathers heavier than a Pound of Lead.



601. If two bodies, one of which is more than the other, be found equiponderant in dinary process of weighing by a balance

larger body is the heavier.

602. Let the bodies in question be those sented within the receiver of an air-pump, annexed figure. On withdrawing the \cdot means of the pump, it will be found th larger body preponderates, though prev counterpoised with accuracy.

603. Rationals.—It appears from a precedia, tration, (573, &c.) that, when any body is sur ed by a fluid, it is buoyed up with a force in p tion to the weight of the fluid, and the qu displaced by the body. Of course, the more it occupies in proportion to its weight, the will its weight be counteracted. In the the two bodies rendered equiponderant in a weight of the larger is most counteracted air. Hence, on exhausting the air from t ceiver, the larger body shows a prepond over the other, equivalent to the superior which the air had afforded it.

604. A similar result may be obtained, if gen be substituted in the receiver for atmoair; because, as its specific gravity to that air is only as I to 14 nearly, each body wor 13-14ths of the support which the air had all

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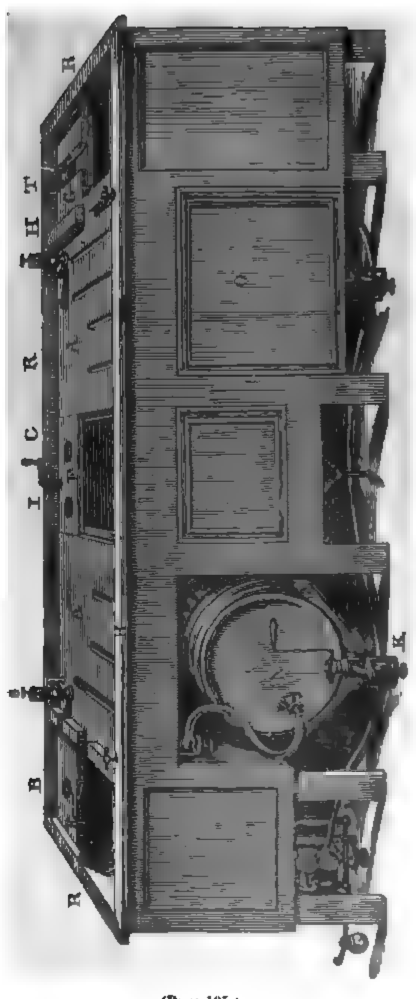
but the larger body, having received more, would lose more. It follows, the common saying, that "a pound of feathers is as heavy as a pound of lead short of the truth; as they would really prove heavier were the air removed.

Table of the Specific Gravities of the Principal Permanent Gases: also of the of 100 Cubic Inches of each Gas.

605. This table is inserted here for convenient reference, not as an object of collectively.

							Specific gravity of 60 degrees-	We 100 inc
Air -	•	•	•	•	•	•	1	31
Oxygen -	•	•	•	•	•	•	1.1111	3
Chlorine -	•	•	•	•	•	•	2.5	71
Protoxide of	chlorine	•	•	•	•	•	2.4444	74
Hydrogen	•	•	-	•	•	•	U 0 6 94	1 1
Steam -	•	-	•	•	•	•	0.625	15
Chlorohydric	(muriatic)	acid	•	•	•	•	1.2472	3
Nitrogen -	•	•	•	•	-	•	0.9722	2
Nitrous oxide	•	•	•	•	•	•	1.5277	41
Nitric oxide	•	•	•	•	•	•	1.04166	3
Ammonia.	•	•	. •	•	•	•	0.59027	11
Sulphurous a	cid -	•	•	•	•	•	2 2222	6
Sulphydric ac		retted	hydrog	en)	•	•	1.1805	3
Carbonic oxid		•	•	•	•	•	0.9722	2
Carbonic acid	1 -	-	•	•	•	•	1.5277	4
Carburetted i	hydrogen (light)	•	-	•	•	0.5555	l i
Olefiant gas	•	•	•	•	•	•	0.9722	9
Cyanogen	•	•	•	•	•	•	1.8055	5
Chloroxycarb	onic acid	•	•	-	•	•	3.4722	110
Fluosilicie ac	id -	•	•	•	•	•	3.6111	lii
Fluoboric aci		•	•	•	·	•	2.3622	1 7

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SECTION IV.

DEFINITION AND DISCOVERY OF THE AERIFORM FLUIDS CALLED GASES.

606. It appears from the na of calorific repulsion, hat solid ponderable mat by combining with caloric, ly passes into that elasist expands, next melts, and te state of fluidity, in which t repulsive power so far redominates over the attr tive, that the particles recede from each other as far a external pressure will permit. When a substance is natu y aëriform, it is called a gas: when it retains the form c r only, in consequence of extracedinary (238,) heat, or removal of pressure, it is caled a vapour.

modified by impurities, until Dr. Black ascertained the nature of carbonic acid gas. Incited by this discovery, oxygra, nitrogen, hydrogen, chloriné, and many other subtraces succeptible of the gaseous state, were discovered, or distinguished, by Scheele, Priestley, Cavendish, and

Others

Of the Art of Collecting and Preserving the Gases.

More Cisterns filed with water or mercury, called hydro-pneumatic or mercuro-pneumatic according to the liquid employed, are used for collecting asses. The vessels intended to contain the gas are filled with water or many, and placed, in an inverted position, on a shelf, or part of the cistrated just below the surface of the liquid. As their orifices are not many shows the surface, they remain full of the liquid, in consequence of the preserve of the atmosphere. (86, 87, 132.) Any gas emitted under the most of a vessel, so filled and situated, rises to the top and displaces the contained liquid.

Hydro-pneumatic Cistern.

In the Appendix will be found an engraving and description of a hydro-pneutic cistera, which I employed in the experimental illustrations of my lectures for that ten years; and which I should probably continue to use now, had not the mand of water from the public works, put it into my power to dispense with the manism for keeping the water at a proper level. As I am now situated, any determined of water is easily supplied from the pipes known here as the hydrant pipes, by the city is supplied with water; and any excess is carried off by a waste pipe. A A rece opposite engraving) is a water-tight platform, surrounded by a wood-ma, R R R R, rising above it about an inch and a half. B, C, T, are three wells with a limit of the form of a hollow parallelopiped, with all of which the cavity leads by the rim communicates; so that when supplied with water to the level of the size of an inch.

GLL L.T. G, are shelves, which severally move in grooves over the wells, so that by may be placed in the most convenient position. Under H is a waste pipe. At

I, is a winch which serves to let in water from the public reservoirs. K, is a 1 for emptying the wells and casks, with all of which, by means of socks, it may made to communicate when requisite. N, is a cask which acts as a gas-holder, ing a communication with the cistern for letting in water from that source; the fices being controlled by valves. By means of a pipe proceeding from its vertex, gas-holder communicates with a pipe or cock, at s, furnished with a gallows and To this, flexible leaden pipes may be attached, for transferring gas either from gas-holder to a bell glass, or from a bell glass to the gas-holder. When a communication is established between the cavities, either of these offices may be perform accordingly as the pressure within the holder is made greater or loss than that of atmosphere. It will be greater when the valve for the admission of water is open that for letting it out being shut; and less when these circumstances are reverses 612. Another cask with pipes and cocks, similar to that represented in the engineer is conscelled by the represented in the engineer is considered in the engineer in the en

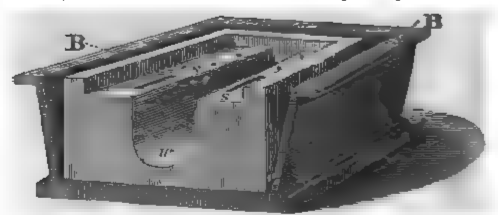
ing, is concealed by the pannel, O.



613. This cut affords a view of the lower side of sliding shelf, in the wood of which will be seen to excavations, T, T, converging into two holes. I shelf is loaded with an ingot of lead at L, to present the converging into two holes. it from floating in the water of the cisters.

Mercurio-proximatic Cistern.

614. The following figure represents the mercurial sistern used in my laborate. The front is supposed to be removed, that the inside may be exposed to view.



615. B B, is a wooden box, which encloses the reservoir so as to catch say of metal which may be propelled over the margin of the cistern. This box is bottom upon stout pieces of scantling, tenanted together and grooved so as to conduct mercury towards one corner, where there is a spout to convey it into a ressel, a ated so as to receive it. The cistern itself is made out of a solid block of w

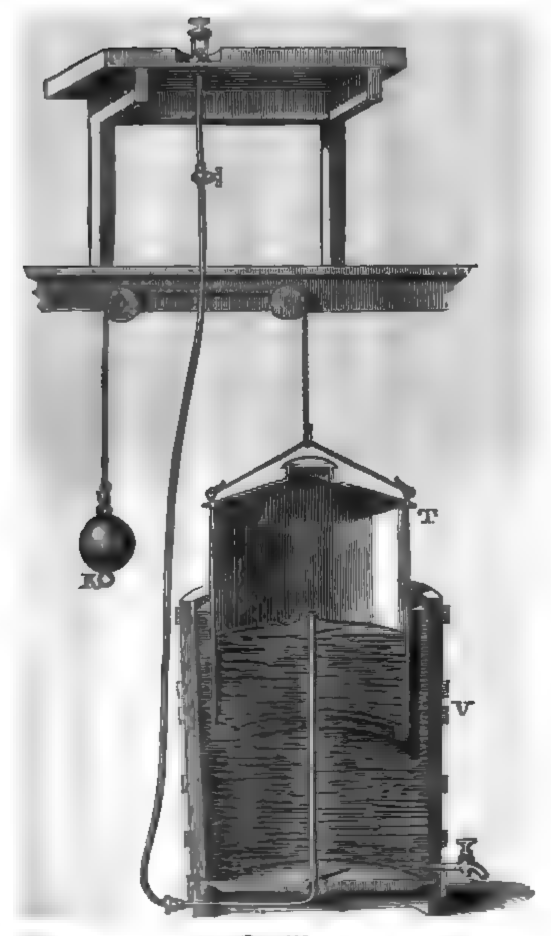
marble. It is 27 inches long, 24 wide, and 10 deep.

616 The ledges, S. S., answer for the same purposes as the shelves in the hyppenmatic distern described in the preceding article. The excavation, w, constitution, w. the well. In this well vessels are filled with mercury, in order to be inverted: placed while full on the ledges. There are some round holes in the markle for troducing upright wires to hold tubes or endiometers; also some oblong mortices allowing the ends of tubes, duly recurved, to be introduced under the edges of t sels to be filled with gas, and in cases of rapid absorption, to afford a passage for mercury into vessels, into which its entrance might be impeded, in consequence their close contact with the marble of the reservoir. To fill this reservoir requi mearly 600 pounds of mercury.

Large Gasometer for Ozygen.

617. The opposite engraving represents a section of my gasometer for eavger other gases, which is capable of holding between eight and nine cubic feet of It is represented as it was situated, when the drawing was made, in the caller at my lecture room. It is now placed in the lecture room in front of my table, one end. The wooden tub, V, is necessarily kept nearly full of water. The c drical vessel, T, of tinned iron, is inverted in the tub, and suspended and compoised by the rope and weight, in such manner as to receive any gas which proceed from the orifice of the pipe in its axis. This pipe, passing by means water-tight juncture through the bottom of the tub, is extended to a cock fixed

Large Gesometer for Oxygen.



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eavity made in the plank forming the rim of the pneumatic cistern. Hence by means of this cock, and a leaden pipe soldered to a brass knob, properly perforated, a communication may be established between the cavity of the gasometer and any other ressel, for the purpose either of introducing or withdrawing gas. In filling this gasometer, the copper vessel and bell glass, used in obtaining nitrous oxide, may be employed advantageously; or the counter-weight being made heavier than the ress. to appending additional weight to the ring, K, the gas may be sucked in from a bell glass, situated over the pneumatic cistern, as fast as it enters the bell from the generating apparatus.

As the gas displaces the water from the cavity of the vessel, T, the latter because were booyant, and consequently rises. When any gas is withdrawn or

Exited. the water resumes its place, and the vessel sinks.

Gammeters which contain 40 or 50,000 cubic feet have been constructed upon principle for holding the gas from oil or coal. They are usually hollow parallel-species. The upper vessel is generally made of varnished sheet iron, the lower one strick work or cast iron. The space within the lower vessel, which is included by the upper one when down, is filled up, so as to lessen the quantity of water required. (See article on carburetted hydrogen.)

INORGANIC CHEMISTRY,

OR CHEMISTRY OF INORGANIC SUBSTANCES.

OF SIMPLE PONDERABLE ELEMENTS, THEIR REACTIONS WITH EACH OTHER, AND THE RESULTING COMBINATIONS.

M. Having in the preceding pages treated of certain general properties of ponderable matter, or those means of ascertaining or observing them of which a knowledge is indispensable to a chemist, I shall, in the next place, proved to the consideration of ponderable substances individually, and their reactive and combinations with each other.

641. In treating of pronderable elements and their multifarious compounds, 127 to arrangements have been pursued by different writers. Some have in the ingin with elements, and to proceed to compounds; others to begin ** - nr ar. is, and to proceed to elements. In favour of the last men-Latin res, it may be alleged, that the most interesting substances in nature her was to us at first, in a state of combination. Thus, for instance, the an enter, salts, acids, alkalies, also flesh, sugar, farina, and other orgraphous, valuable either as food or as medicine, are compounds which have been naturally made the subjects of chemical inquiry; and it may be The student might with advantage be induced to travel in those parts, of which a successful pursuit has led to that chemical knowledge The state object to impart. In this way he proceeds from facts which * are well to such as he ought to learn, in the order in which he would promoter's advance as far as he might be competent. But it may be thered, that no somer are the ingredients of a body stated, than the stube a distracted by names, of which he is ignorant; and which there is an mediate percessive to explain. Hence it follows that the ingredients of a care and may come to be considered in immediate succession, when they have no analogy with each other; while it is highly advantageous, after having treated of any one element, to proceed to that which has the greater analogy with it. In that case, a certain portion of the conceptions which have been formed respecting one element, may be extended to another, while mental exertion, and without much additional pressure upon the memory.

stance first, and afterwards of compounds, is objectionable, because it can be put into practice effectually. To treat of the chemical habitudes o one element, requires that we should speak of other elements, in remains with which, those habitudes are displayed, and respecting which a beggins of course ignorant. In pursuing this course, each substance must reated of imperfectly, or language and illustrations employed, which student is unprepared to understand.

623. The course which I have chosen is as follows. I begin with the ment which, of all ponderable matter, has the most important part assist to it in nature; I mean oxygen. The history, state of existence in and means of procuring, and properties of this substance, so far as they can rendered intelligible to a novice, are stated, or exemplified and explain In the next place to oxygen, I present chlorine to attention, which has least as much analogy with oxygen, as any other known element, and at the same time, an agent of high importance. Having treated separat of oxygen and chlorine, as far as may be expedient, the compounds with they form with each other, may, in the next place, to a certain extent, treated of with advantage. Then, guided by analogy, bromine and iod though inferior in importance, may be successively treated of, and su quently all the compounds which they can form, either with oxyges or c rine, or with each other. This system will be followed in treating of the elements.

624. Pursuant to this method, little can be said of fluorine in the sec appropriated to its consideration, since those elements with which its t interesting reactions take place, cannot consistently be made the object attention under that section.

625. Cyanogen is, in its properties, analogous to chlorine, brown and iodine, yet being composed of carbon and nitrogen, should not be object of attention, until the pupil is prepared by a knowledge of its constituents. Besides, it comes in consistently under the general heat carbon, which, agreeably to my plan, as above explained, comprises compounds of carbon with all substances previously treated of, among w is nitrogen.

626. Of the fifty-four simple elements universally recognised by mists, a list, with their equivalent numbers and symbols, has already

given. (560, &c.)

627. Of these elements, chlorine, bromine, iodine and fluorine are cla by Berzelius under the name of halogen bodies, or generators of salts; v oxygen, sulphur, selenium, and tellurium are classed together under name of amphigen bodies, or both producers; meaning that they are ductive both of acids and bases. To the elementary halogen bodies adds the compound body cyanogen. I object to this classification, tha word salt admits of no definition, reconcilable with the use which has made of it by the distinguished author; and because, from facts and nitions practically sanctioned by him, and chemists in general, it is evi that the elements belonging to both of his classes are productive of and bases. Hence I have associated them in one class, under the app tion of basacigen elements. In honour of Berzelius, I shall, however tain the terms halogen and amphigen, in order to designate the elen which he has distinguished by those names. It may be proper to add we owe to Berzelius himself the idea that any other substance besides gen could form acids and bases capable of uniting to form salts.

swiege of the existence of this faculty in three of his amphigen elests, sulphur, selenium and tellurium, is, I believe, entirely due to his selenium. If chemists, myself among others, who consider his double to account ing of acids and bases, are in the right, it is to the light aftial by his brilliant discoveries that we owe the ability to pursue the true

Before concluding this preliminary exposition of the classification amendature which I propose to adopt, I wish to make it clear, that which of producing both acids and bases, which, agreeably to the servelius, is restricted to his four amphigen elements, is, agreeably inc, extended to the elements comprised in both of his classes, which consequently united under one designation, as basacigen elements. My cigen class is, therefore, the amphigen class of Berzelius, enlarged rance and more descriptive name, * so as to take in both of his paradamphigen classes.

Is order to render the definition of a basacigen body precise, it recessary that I should give a definition of acidity and basidity.

I shall proceed to give a definition which to me appears quite tory. It is perhaps necessary to premise, that a tertium quid recally to the old chemists, a compound in which the qualities of edients were neutralized, or so much altered, as to make a body of a chemical reaction differing from that of either of its ingredients. It is as follows.

But to proceed finition; it is as follows.

When of two compounds capable of combining together to form a paid, and having an ingredient common to both, one prefers the the other the negative pole of the voltaic series, we must deem the maid, the latter a base.‡

Thus sulphuric acid, (consisting of sulphur in combination with and soda, (consisting of sodium and oxygen,) are capable of comform sulphate of soda, a tertium quid. Each of these compounds ommon ingredient, oxygen, and one of them, the acid, prefers the sole of the voltaic series, the other the negative pole. It follows, huric acid is entitled to the appellation of an acid, while soda may at of a base.

I, I trust, be perceived, that a basacigen element is one capable of proth an acid and a base, the monosyllable gen being understood, in chemical, when added to a word expressive of a property, or state, to signify the producing that property, or state. (633.)

t mane is much needed to convey the idea of the basic property, as set the acid property, I have ventured, without any authority, to embedding, which from its analogy with acidity, must, I presume, be pinalligible.

it to be understood. that I consider this definition as only declaratory of the chemists, who all obey the rule, although, as far as I know, excepting it has never been enunciated.

t doesn it necessary to introduce into the text a corollary, which inless from the cited definition, as it would unnecessarily distract attenit may be well before taking leave of this subject, to say, that agreeably all gractice, any body which is capable of saturating a base, is considered is said that on the other hand, any body which is capable of saturating an allowed to be a base. It is upon this basis that the pretensions of the orlies and acids to be considered as acids or bases, are founded.

OF INDIVIDUAL PONDERABLE ELEMENTS,

AND OF THEIR REACTION WITH EACH OTHER, AND TERESULTING COMPOUNDS.

ed, (627,) eight being designated as basacigen, make, with cyanogen, the compound basacigen body, (629,) nine in all in the basacigen class. I shall designate the rest of the elements as radicals; subdividing them into metallic radicals and non-metallic radicals.

OF BASACIGEN ELEMENTS.

634. Oxygen,
Chlorine,
Bromine,
Iodine,
Fluorine,

Cyanogen, Sulphur, Selenium, Tellurium.

They will be treated of in the order in which they have

been named, in the eight following sections.

should employ his appellations amphigen and halogon. There is, in fact, a necessity for words to distinguish the bodies to which he has applied these names; especially from the very great analogy between those which are dissignated as halogen.

636. The student is requested to recollect that chlorist bromine, iodine, fluorine and cyanogen constitute the lightness of Berzelius, while oxygen, sulphur, selenist

and tellurium form his amphigen class.

SECTION I.

OF OXYGEN.

637. In the gaseous state, oxygen forms one-fifth of atmosphere in bulk; and as a constituent of water in ratio of eight parts in nine, it pervades every part of creation where that important compound is to be for It exists in that congeries of oxidized matter which call earth, and is a principal and universal constituent

al and vegetable matter. Its combinations with meand various other combustibles are of the highest tance in the arts. It was called oxygen under the cous impression of its being the sole acidifying prinfrom the Greek ozo, acid, and yrropes to generate.

. Preparation.—It can only be isolated in the form as. It is yielded by red lead, nitre, or black oxide nganese, when exposed to a bright red heat in an ottle. There are various other means of obtaining n gas. It is generally supposed that, in order to it in a high degree of purity, chlorate of potash be employed; but I have found the first portions of is as evolved by a red heat from nitrate of potash or e of soda very nearly pure; and Dr. Thomson althat this salt, by exposure to a carefully regulated parts with one-fifth of the oxygen of its acid in a of purity; or in other words, it gives up an atom of in for every atom of the salt, which is equal to 8 in 102 parts, or rather more than one-thirteenth.

9. Properties of Oxygen.—The specific gravity of this atmospheric air being 1., is 1.1024. One hundred cunches of it weigh 34.1872 grains. In refracting light, we is inferior in power to any of the other gases.

but slightly absorbed by water, does not differ from non air in appearance, but is somewhat heavier, and orts life and combustion more actively. Under a bell belief with oxygen gas, an animal lives, or a candle thrice as long, as when similarly situated with the equantity of common air. Oxygen gas is supposed onsist of oxygen rendered aëriform by caloric. The ivalent of oxygen is 8, hydrogen being unity.

Apparatus for obtaining Orngen upon a large Scale.

As nearly as much time and trouble are expended in conducting a chemical was a small scale as upon a large one; and as in my experiments I consume parties of oxygen gas, I have lately employed the cast iron alembic repressive following figure, for the purpose of obtaining the gas from 12 or 15 a.m. When in operation, it is made to occupy a suitable cavity in a large. When in operation, it is made to occupy a suitable cavity in a large. The neck is so formed as to receive a large hollow knob of iron, from harrel proceeds at right angles. This knob is secured by a gallows have a saturated solution of borax. To the orifice of the gun barrel, the pipe is attached, by which the gas is conveyed to the gas-holders

Constaken to use no more fire than will bring over the gas, and the opera-



precautions, the gas is of better quality; the first portion being mearly pure, and the alembic is less correded. Besides, the nitrate, being converted into nitrits of potasis produces, by deflagration with charcoal, a telerably pure carbonate of potasis.

Experimental Illustrations of the Properties of Oxygen Ga

- 643. Several cylindrical glass vessels of an appropriate shape being filled with the gas over one of the pelves of the pneumatic apparatus, the following illustrations of the energy of oxygen gas in supporting combustics are alforded.
- 644. Let a stout wire be made, at one of its ear, to en brace a lighted candle, so that it may be convenient lowered into the bell while replete with oxygen. It wis be found that, if the flame be extinguished, and the considered into the gas while the snuff remains red-lot inflammation will be renewed with great energy.
- 645. The vessel being replenished with gas, the fame? a piece of burning caoutchouc, let down into it, acquired dazzling brightness.
- 646. Let Homberg's pyrophorus, or preserably such and have contrived to obtain from Prussian blue, or tanno go late of iron fall through the gas. During its descent to pyrophorus takes fire spontaneously, producing an igner shower.

47. An analogous fiery shower results, when charcoal der, or filings of iron or steel, made red-hot in a cruciare projected in like manner into oxygen.

18. If (by means of a blowpipe,) a jet of oxygen be to act upon a lamp flame, or upon that of hydrogen, her pure or carburetted, an intense heat will be ext. (379.)

19. An iron wire, being heated in the flame thus exl by oxygen, takes fire, and continues to burn spleny, although the lamp be removed.

Combustion of Iron or Steel in Ozygen.



pipe communicating with a cock of one of the air holders supplied with oxygen, a glass vessel, such as is usually employed to shelter candles from currents of air. Let the upper opening of the vessel be closed by a lid with a central circular aperture, as represented in the engraving. Leaving this aperture open, by turning the key of the cock, allow the gas to rise into the vessel, from the holder. Next apply a taper to the aperture, and as soon as it indicates, by an increased brilliancy of combustion, that oxygen has taken the place of the air previously in the vessel, cover the aperture. In the next place, attach a small piece of spunk to one end of a watch spring, or of a spiral wire as in the figure. Ignite the spunk, and removing the cover, plunge the end of the spring associated with the spunk into the gas. The access of the oxygen causes the spunk to be ignited so vividly, that the spring or wire, takes fire and burns with great splendour, forming a brilliant liquid globule, which semullates beautifully. This globule is so intensely hot, that sometimes, on falling, it

momentally sink into the water, but leaps about on the surface, in consect the steam which it causes the water to emit. If it be thrown against the I the containing vessel, it usually fuses it without causing a fracture, and has hown to pass through the glass, producing a perforation without any other. These phenomena are more likely to be produced when an iron wire is this experiment, then when a steel spring is employed, as the fusing point of the tree is higher than that of steel.

Necessity of Oxygen to a Cundle Flame demonstrated.

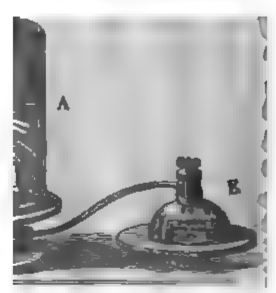
L Accords will burn only for a limited time in a limited supply of air, it will weren, but burns brilliantly in oxygen gas, and much longer than in a process of air.

14

C

B

D



first there be two bell glasses, A and B, communicating with each ether fiexible leaden pipe, a cock intervening at C. Suppose A to be placed over high candle on the plate, D, which communicates with an air pump plate as appear at E. It will be found that the candle will gradually burn more dimly, and wi last go out, if no supply of fresh air be allowed to enter the containing bill. It repeating the experiment, the air be withdrawn by means of the pump, the each rapidly extinguished. 'It is thus demonstrated that a candle will not been in we and that it can burn only for a limited time, in a limited portion of atmospheric and that it can burn only for a limited time, in a limited portion of atmospheric with the receiver, B, filled with oxygen over the pneumatic cistern, the water rise into and fill the receiver, while the gas will be transferred to the bell, means of the galvano-ignition apparatus (335), the candle may be again lights the oxygen, when it will burn splendidly.

Combustion of Phosphorus in Ozygen Gas.



654. A brass plate, which answers as an extra air pump plate, is supported bollow cylinder of the same metal. Concentric with the axis of this cylinder

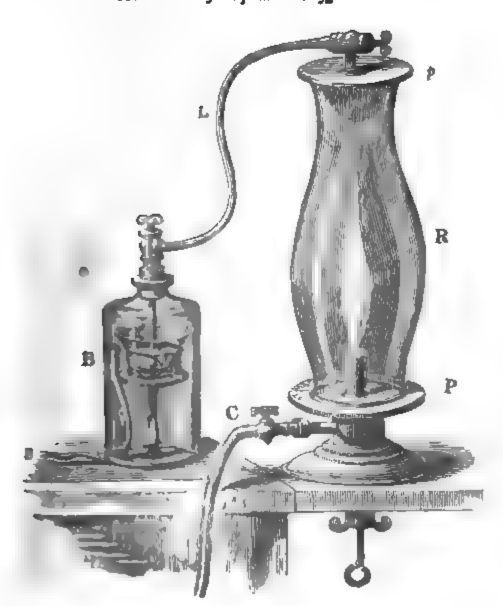
OXYGEN: 115

exthrough it. so as to reach about three inches above the plate, there is a cost three-fourths of an inch in diameter, open below, but closed above by copper disk to which it is hard soldered. The tube is fastened into the or a brass screw plug, in the centre of which the tube is soldered. Hence, the bore of the tube is accessible from below, so far up as the concave copsing aurinounts it, no air can pass through it, or through the cylinder. bout twenty grains of phospherus being placed upon the copper disk, are is put over it upon the plate; and by causing one of the pipes which are laterally to the cylinder to communicate with an air pump in operation, the mainsted. By means of the other pipe, a due quantity of oxygen gas in from the bell glass, B, to which this pipe is annexed. The apparatus is prepared, the end of an iron rod previously reddened in the fire, is passed the bare of the tube so us to touch the copper disk which holds the phos-The most vivid ignition ensues. The light has at first a dazzling beauty, as shore of its beams" by the dense white fumes of phosphoric sold, which ustion evolves. Hence, an effulgance, approaching to solar brilliancy, soon a milder illumination like that of the moon, rendered more pleasing by the

he gishes with which I am accustomed to perform this experiment contain gallons. It is better that the gas in the globe should be in some degree otherwise the expansion at first excites a considerable effort in the air to The enlargement of bulk, arising from the heat, may be provided for by a adder, a communication with which being opened, a portion of the heated abled to retire, till the condensation of the exygen with the phosphorus, phonic acid, compensates the expansion.

have performed this experiment, when the density of the gas was one-balf if in equilibric with the atmospheric pressure. This of course obviated the y of any all consequences from expansion.

Combustion of Sulphur in Oxygen Gas.



658. Supposing the junctures made by the plates, P p, with the receiver, R, to air-tight, and that there is a communication between it and the bell glass, B, means of a flexible leaden pipe, L, it must follow that, whenever the suction pus from which the recurved pipe, S, terminating within the bell, proceeds, is ma act, the air in B being rarefied, that in R will force its way through L, and the liqu in the vase upon the stand. It must also be evident that, if the pipe and cook, communicate, on one side with the receiver, on the other with a reservoir of exygi this gas will be impelled into the receiver, as soon as the cock is opened, in ec restore the equilibrium destroyed by the suction pump.

659. The plate, P, with its supporting hollow brass cylinder, has been already scribed in the preceding article. The tube, surmounted by the disk, used in combustion of phosphorus, is removed, and its place a piece of a gun bassel i like manner, fastened, so that the butt-elm may occupy the axis of the ex The touch-hole being closed, a perforation. similar in size, is drilled in the the barrel, at the point from which the figure. In order to produce this jet of about the end of a rod, and tied on i... brimstone. The gun barrel, during a t butt-end, where it is perforated. Being we wed into its place again, the sed, with the cotton and sulphur, is pushed up to the bore of the burrel. By the of the iron, the sulphur is converted into a . ot vapour, which, issuing in a jut

ed sulphur, some cetton wick is s tuft, thus made, in souked in cary removal, is heated red hat the perforation, enters into combustion with he oxygen in the receiver.

660. In consequence of the rarefaction wi the air in the bell, B, by the pump, the fumes of the burning vaporized a liphur are drawn through the the vase upon the stand, in which, consequently, a mixed solution of sulphusous solution is such as a sulphusous solution in sulphusous solution is sulphusous solution.

sulphurous acids is produced.

Additional Illustration of the Combustion of Sulph in Oxygen.

661. The preceding illustration has not for two three years, been exhibited before my class, yet presumi it might not be uninteresting to some of the students this work, I have not omitted it from this edition. terly I have resorted to the following method of exhibiting the combustion of sulphur in oxygen, as being easier, as yet sufficiently pleasing and instructive. All the steps the process for the combustion of phosphorus in oxyge are performed, as already described (651, &c.) but in li of a stick of phosphorus, a tuft of asbestos soaked; melted sulphur, is placed upon the capsule, with a minu piece of phosphorus beneath it. The latter when heat by the incandescent iron takes fire, and consequently nites the sulphur, with which the asbestos is imbued. whiteness and dazzling brilliancy, the light afforded by t combustion of sulphur in oxygen is inferior to that evolve by phosphorus, when similarly situated; but this inferiori is compensated by the splendour of its characteristic pu ple hue.

SECTION II.

OF CHLORINE.

. As a gas, chlorine exi y by artificial means; an ingredient in marine t, in the proportion of ifths, it constitutes nearly fiftieth of the matter ocean, and is widely dis nated throughout the an ingredient in some s well as the sea. It is most active agents used in chemistry or medicine. discovered by Scheele, and called by him dephlogis-I marine acid. It afterwards received the name of nated muriatic acid, or oxymuriatic acid, from Laand the chemists who adopted his nomenclature, the erroneous idea that it was composed of muriatic ed oxygen. Its present name was given by Sir H. from zage green, because its colour is greenish.

Preparation.—It is obtained by heating in a retort mbic, of glass or lead, three parts of black oxide of mese, with four parts of muriatic acid; or the same ty of this exide, with eight parts of common salt, four

of suphunc acid, and four parts of water.

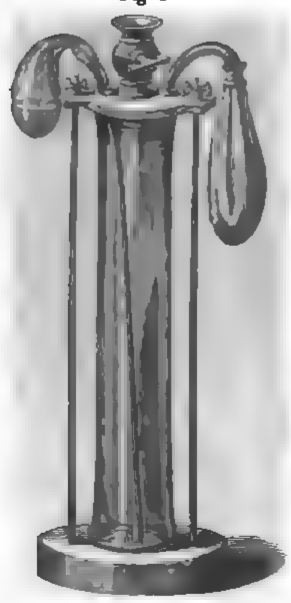
L. Being a gas, chlorine must be received over the pneumatic cistern in bell glasses or bottles; the temre of the water should be raised, by adding a porolling hot. As much of it is absorbed if it remain a contact with the water, I generally employ glass with air-tight stopples, in order that they may be red from the water as soon as filled. Berzelius althat if the water employed be saturated with salt, is less absorption.

Lars or bottles may be filled with chlorine gas, by of a tube or retort beak, as in fig. 1, of the follow-praying, reaching from the generating vessel to the nof that into which it is to be introduced. The air placed by the chlorine, in consequence of its superior by, without any admixture ensuing adequate to intertable the exhibition of its characteristic properties.

When substances which take fire in the gas are to bedeced, it is expedient that a communication should the inside of a bladder attached, as in the followers, which represent apparatus, of which fig. 1 med for the combustion of metallic powders, fig.

2 for that of phosphorus, introduced by means of the dle L.









667. Properties.—When pure and dry, chlorine is a manent gas of a greenish-yellow colour. Its weighthat of common air, is nearly as two and a half to Even when existing in the air in very small proportic is intolerable to the organs of respiration, and to respipure, would quickly produce fatal consequences.

668. Mr. Faraday has shown that, under great preschlorine becomes a liquid. It will remain liquid sometants after all pressure is removed, in consequence of

great cold produced by its evaporation.

669. That species of chemical action which is atte with the phenomena of combustion, is supported by gas with great energy. It combines directly with e combustible except carbon. It has a curious property, noticed by me I believe, of exciting a sensation of war though a thermometer, immersed in it at the same

that its temperature is greater than that be adjoining medium. The heat thus noticed is probated by a reaction with the matter insensibly per-

- 0. Chlorine is absorbed by water, and the solution powerfully on metals. It appears to be the only solof gold. At the temperature of 40°, it forms with a solid hydrate, consisting of 1 atom of chlorine,) atoms of water. Silver, in solution, is the best test orine; and, reciprocally, chlorine is the best test for red silver. The compounds of chlorine with merso useful in medicine, will be treated of when on the t of that metal. When the aqueous solution of ne is exposed to the solar rays, it forms muriatic rith the hydrogen of the water, while the oxygen s. It bleaches by liberating the oxygen of water, hus enabling it to act on the colouring matter. igh it has no direct reaction with oxygen, when in ascent state, these elements unite to form four coms, all of which are now considered as acids.
- About thirty years ago, chlorine gas was univerconsidered as a compound of muriatic acid and oxyund called oxymuriatic acid. It is now deemed an utary substance, rendered gaseous by caloric.

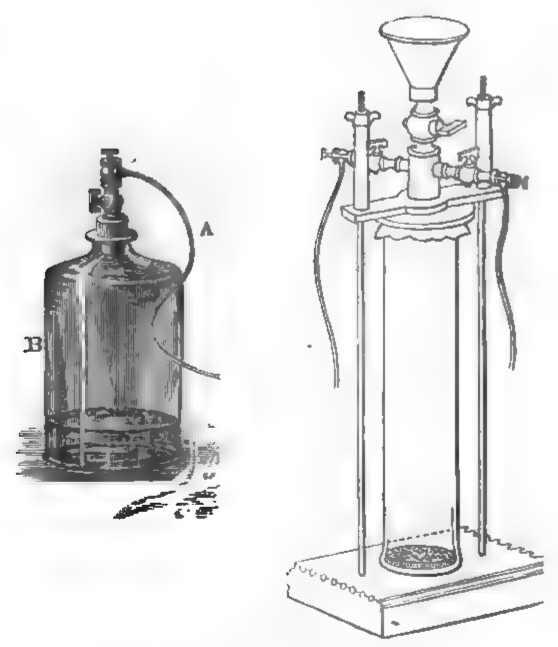
xrimental Illustrations of the Properties of Chlorine.

- Leaves of Dutch gold, introduced by means of a rod into a bottle of chlorine, take fire.
- 3. Calorific influence upon the fingers compared with upon a thermometer.
- 4. An infusion of litmus whitened in descending in a n through the gas from a funnel.
- i. A lighted candle introduced, burns with a carboas flame.

Combustion of Intimony in Chlorine.

Then an air pump is at hand, the following apparatus may be used for the im of powdered antimony. It consists of a large jar closed air-tight, and in the screw rod and plate frame described. (24%) by means of one of the flexible pipes and cocks with which the apparatus is a communication may be made with an air pump, and with a large vessel, making chlorine. Into the centre of the lid a cock is fastened, the key of means of being perforated as usual, is drilled only half through, so as to propose a capable of holding a thimbleful of powder. The cavity in the bessel is charged with pulverized antimony, which, on turning the key half through the chlorine, and as it falls assumes the appearance of a shower

of fire. The cock being, from its construction, always closed, and the being tight, the spectators are protected from the nozious fumes.



678. In this experiment, the chlorine forms with the antimony a compound has less capacity for caloric and light than its ingredients have separately by their combination, the phenomena of combustion are produced. The patho combustion is the perchloride.

Apparatus for the Combustion of Metallic Leaf* in Chlorine.

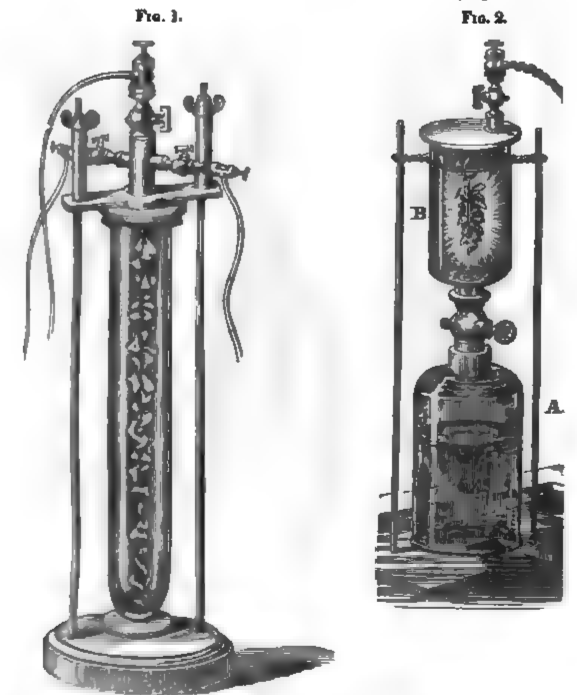
67%. The apparatus used in this experiment (See fig. 1,) differs but little one above represented, (677, &c.) being the same as that described in (241, &c.) excepting the funnel, which is unnecessary in this case.

680. Into the lower end of the cock a rod of iron is screwed fast. This

680. Into the lower end of the cock a rod of iron is screwed fast. This such dimensions as to extend from the top to the bottom of the receiver, a ported within it, so as to be in its axis or every where equidistant from the Before fastening the plate into the situation in which it is represented in the it must be lifted in order to attach the leaf metal to the rod with the air arabic. The arrangements being so far completed, the cylindrical receive been exhausted by means of the air pump, the cock, regulating the comme with that instrument, is to be closed, and the other which controls the enthe gas is to be opened. By these means the leaves burn splendidly, being neously enveloped in an atmosphere of chlorine, which rushes in to supplement caused by the air pump.

[&]quot;The metal usually employed is the Dutch gold leaf of the shops, an a cipally of copper and zinc.

63. Anther method of performing this experiment is illustrated by fig. 2.



The metallic leaves being suspended from the plate which closes the bell, B, d the bell being exhausted of air by means of the pump, chlorine is suddenly adted into a by the glass cock from the bell glass, A, previously supplied with the

Spentaneous Combustion of Phosphorus in Chlorine.

The figure at the top of the next page, is intended to convey an idea of the standard inflammation of phosphorus in chlorine, by means of an apparatus which the lecturer to perform the experiment without exposing spectators to the ma. Let there be a cylindrical glass vessel, eight or nine inches in diameter, about a foot in height, with a neck about four inches high, and one and a linches in bore, the whole resembling a large decanter without a bottom, set the orifice of the neck, let there be cemented, air tight, a brass cap, surmed by a stuffing box, and having on one side a hole communicating with early of the neck. This aperture must be furnished with a screw, by which it is spended or closed at pleasure. Through the stuffing box a copper rod passes, to been end of which a glass or leaden stopple is so affixed, as to close the lowest of the neck, into which it is ground to fit air-tight. Over this stopple, a cup there is soldered, so as to be concentric with the rod. The rod terminates above i smalls. Within the cup, let ten or fifteen grains of phosphorus be placed. It smalls the fitting that the cup, let ten or fifteen grains of phosphorus be placed.



the cavity of the vessel, by a suitable movement of the sliding rod. In the place draw up the cup and plug into the neck, so as nearly but not entirely int, and sink the vessel into the water of the pneumatic custern until all the authorized in the screw, and the plug twisted and drawn into its place, so as to tight. After filling the body of the vessel thus with water, place it upon the the cistern. Chlorine may now be allowed to occupy three-fourths of the within the vessel below the plug. The process being so far advanced, it is occasary, at the moment when it is desirable to produce the combustion, to the plug, and of course the cup associated with it containing the phosphor the cavity supplied with the chlorine. The physich russion burns actively, a with a feeble light. The increased temperature consequent to the combicauses the gas to expand, but not so much as to become two bicks, to be retained. In this case the chlorine forms a chloride of phosphorus, which, meeting water, is decomposed into phosphoric and in matical as ds. By transferring the after it is supplied with chlorine, to a clean porcelain or glass dish, covere pure water, the products of this combustion might be saved, and would of coveresse in proportion to the quantity of phosphorus and chlorine employed larger scale, this process might be resorted to advantage such for the gener phosphoric acid, which is produced when the proportion of ch. The is sufficie four cobic inches for every grain of phosphorus.

Of the Compounds of Chlorine with Oxygen, and of the Nomencla these Compounds and others formed with the Basacigen Class

695. Consistently with the French neroenclature, the combinations by oxygen, chlorine, bromine, indine, and distinct, with other elebave been distinguished as acids, or characterized by a termina "ide," or in "ure," which last monosyllable, when there has been tention of altering the meaning, has, by the British chemists, been tra-

and wret. The termination in ide, which is common to both languages, is, by Theoard, and other eminent French authors, restricted to the binary comreads of oxygen which are not acid. Analogous compounds formed with the halogen elements, chlorine, bromine, iodine, fluorine, cyanogen, &c., have by the same writer been designated by the termination in ure. we have in his work, chlorures, bromures, iodures, fluorures, and cyanures. Some of the most eminent chemists in Great Britain have distinguished the elements called halogen by Berzelius, together with oxygen, as supprovis of combustion, and have designated the binary compounds made with them, when not acid, by the same termination as the analogous commands of oxygen. Accordingly, in their writings, instead of the names nentioned, we have chlorides, bromides, iodides, fluorides. 7 s Chamstry, cyanure is represented by cyanide; in Thomson's, by cyanwinder: and in Brande's and Turner's, by cyanuret. I shall follow the practhe of the British chemists in the case of the four first mentioned compunk, extending it to the compounds of cyanogen, as Henry has done.

These rules of nomenclature will be considered as extending to all the isoligen class. Of course, the compounds of sulphur, selenium and the isoligen, when not acid, will be designated by appellations terminating in ide. In lieu, therefore, of sulphuret, selenuret and telluret, I shall in common with Berzelius, employ the words sulphide, selenide and telluride.

COMPOUNDS OF CHLORINE WITH OXYGEN.

687. 1 atom or 1
volume of chlorine
volume of chlorine
volume 36. forms

With 1 atom, or 3 volume of oxygen, or hypochlorous acid, 44
With 4 atoms, or 2 volumes of oxygen, chlorous acid, 68
With 5 atoms, or 23 volumes of oxygen, chloric acid 76
With 7 atoms, or 33 volumes of oxygen, perchloric, or oxychloric acid, 92

Of Hypochlorous Acid.

This compound, of which the ingredients are stated above, (687,) represented by the reaction of chlorine, with an excess of finely pulverized proximal of mercury, suspended in pure water by agitation.

123. &c..) combines with both the oxygen and mercury, forming two companies, a bichloride of mercury, and a protoxide of chlorine, or hypochloms acid, which dissolves in the water. The bichloride combines with a potent of the undecomposed bioxide, and forms a kind of combination, generally designated as oxychloride, indicating that a substance so called consists of an oxide, and a chloride. The oxychloride formed in this case, bing almost insoluble, is separated by filtration. A more concentrated solution of the acid is procured by successive distillations, in which as little had as possible should be used, and preferably it should be accomplished by successive pressure.

Properties of Hypochlorous Acid.—The aqueous solution of hypochlorous acid, resulting from the above described process, is, when concentrated, in colour slightly yellow, with an odour strong and penetrating,

For the purpose of distillation, by reduced pressure, the apparatus represented and described in page 69, (39%) might be used, substituting a second and third retert, well refrigerated, for the vessel, B, and the bottle, C. (188.)

resembling that of chlorine, but yet differing therefrom perceptibly; upon : the skin its effects are similar to those of nitric acid, but more active. Its bleaching powers are eminently great. It is so much prone to decomposition, as to undergo that process spontaneously at ordinary temperatures, being resolved into chlorine and chloric acid. This change is accelerated by light, and ensues immediately from direct exposure to the solar rays. Bodies full of sharp corners, (the fragments of powdered glass, for instance,) when thrown into the liquid acid, are productive of an evolution of chlorine with brisk effervescence. The oxydizing powers of this reagent are powerful but various, being most active with non-metallic elementary radicals, such as sulphur, phosphorus, and selenium. Each of these it readily saturates with oxygen, and likewise iodine, and bromine, which are thus severally converted into bromic and iodic acid. Its reaction with gold and platinum, is but feeble, but with iron and silver energetic. The former is converted into an oxide, the latter into a chloride, while in the case of the one, the oxygen escapes, in that of the other the chlorine. Mercury it converts into an oxide, and a chloride, which form an oxychloride, by uniting in their nascent state.

Of Gaseous Hypochlorous Acid.

691. Balard, to whom we are indebted for our knowledge of the facts above stated, was successful in procuring hypochlorous acid in the gaseous form, by introducing into a concentrated aqueous solution over mercury, anhydrous nitrate of lime in successive portions. By its superior affinity for water, the nitrate causes the evolution of the acid in the aëriform state, the mercury being protected by the interposed solution of the nitrate.

692. Properties of Gaseous Hypochlorous Acid.—The gaseous hypochlorous acid much resembles chlorine, in possessing a greenish yellow

colour. Water absorbs 100 times its own volume of it.

693. A slight increase of temperature is sufficient to cause hypochlorous acid to detonate, and though less explosive than chlorous acid, it is apt to explode, when an effort is made to transfer it from one bell glass to another.

694. Its composition was ascertained by Balard, by analyzing the gascous product resulting from its explosion, by which it was found to consist of one volume of chlorine, and half a volume of oxygen, as already stated. (638.)

Of Euchlorine or Impure Chlorous Acid.

695. In the last edition of this work, euchlorine was treated as a protoxide of chlorine, but it was, at the same time mentioned, that doubts existed whether the gaseous substance known by this appellation, might not be a mixture of chlorous acid with chlorine. These doubts appear to have been succeeded by an affirmative conviction, and accordingly I have omitted the name from the list above given, of the definite compounds of chloring with assume

rine with oxygen.

696. Euchlorine is obtained by heating gently, in a small glass retort, equal parts of strong muriatic acid, water, and chlorate of potash. The retort should only be subjected to the flame of a small spirit lamp, or an inflamed jet of hydrogen, which should be so situated as not to heat the body of the retort above the part containing the liquid; as this may cause an explosion. It is advantageous to interpose, as a support for the retort, a plate of tin, having a circular aperture of about an inch and a half in diameter. By these means, the application of the heat may be sufficiently restricted.

1800 The gas may be received over mercury, although not without 1800 nearence; since by its decomposition, in consequence of the large proportion of free chlorine with which it is associated, the mercury is superficially converted into a subchloride. But, while the covering thus formed, protects the surface of the metal from further erasion, it also, by coating the internal surface of the glass, hides, more or less, the remarkably deep pre-nish-yellow colour of the gas from the eye of the spectator.

through a tube, replete with protochloride of mercury, (calomel,) this chorde absorbs an additional atom of chlorine, and thus brings the chlores acid to a state of purity. The rationale of the evolution of the mixture known as euchlorine, seems to be as follows. By double elective affinity, there is a reciprocal decomposition of the potassa and chlorohydric acid, causing the separation of the chloric acid, containing five atoms of oxygen. Consequently, by the reaction with these atoms, a further dehydrogination of chlorine ensues, causing a portion to be set free, while another portion retains enough oxygen to constitute chlorous acid.

Process for claborating pure Chlorous Acid directly.

14 parts of concentrated sulphuric acid, and receiving the gaseous product over mercury. The evolution of the gas takes place without heat at first, but to be completed requires a temperature near to 140°. This should not be exceeded, and the heat should be restricted to the bottom of the retort, as in the case of euchlorine. The process is replete with danger, as from slight causes this gas explodes with surprising force.

Retionale.—By the action of sulphuric acid on chlorate of potash, two compounds are produced, chlorous and oxychloric acid. The former is traced in the gaseous state, the latter remains in union with the potash livers is as if one portion of the chloric acid were displaced from its area with the potash by the superior affinity of the sulphuric acid, and we reinquished a part of its oxygen to another portion of the same acid, so a union with the alkali. The chlorate of potash is thus partially constraint into an oxychlorate. The deoxidized chloric acid constitutes a formula which is designated by Berzelius as chlorous acid. By others, there was variously designated as the tritoxide, quadroxide, or peroxide of cherge, as consonance with the different impressions entertained of its presentes, or composition.

Properties of pure Chlorous Acid.

701. Cherrons acid gas has a yellow colour, which is deeper than that of chorus. Its odour is somewhat aromatic, and bears no resemblance what of choruse. It whitens a solution of litinus, without reddening it. When subjected to an electric spark, or to a temperature of 212°, it expects with great violence, giving out light and heat, and being converted attachoruse and oxygen. Agitating the gas with mercury will sometimes produce the same result. Water absorbs seven times its volume of chlorous and gas, acquiring a deep yellow colour, and a peculiar acrid taste, which is nevertheless not at all acid. The aqueous solution, when added in small quantities, presented to the sour's rays evolves chlorine, while oxychloric acid remains in section. In a diffuse light it takes several weeks to effect this decomposition, and it does not take place at all in the dark. Faraday has found

that chlorous acid gas may be liquefied by subjecting it to great pre The resulting liquid is of a yellow colour.

A convenient and safe Method of effecting the Explanion of Buchlerine.



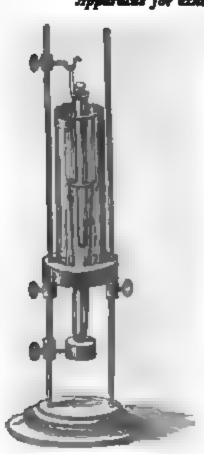
702. A convenient and safe method of effecting the explosion of suchlarine is represented in the preceding figure. The gas being introduced into a strot of about \$\frac{2}{3}\text{ths of an inch in diameter, and fifteen inches in length, over more applying a heated metallic ring, an explosion enques. The gas at the san loses its greenish-yellow colour, and increases in bulk. The chlorine is subset absorbed by the mercury.

703. Thenard advises the application of a spirit lamp to produce the nest temperature. It is easier and more safe to use the hot ring. The tube is of sity supported by an iron wire, which has been overlooked in sketching \$\frac{2}{3}\text{in \$\frac{2}{3}\$}\$ and \$\frac{2}{3}\text{overlooked in sketching \$\frac{2}{3}\text{in \$\frac{2}{3}\text{overlooked in sketching \$\frac{2}{3}\text{in \$\frac{2}{3}\text{overlooked in \$\f

power to caloric, there ought, after an evolution of heat, to be a reduction of in any gaseous compound; but by the decomposition of suchlorine, although is evolved with explosive violence, the volume of the gaseous matter is intre-705. The only explanation which I can give, is, that the capacity for caloric

compound in this case, as in others, is greater than the sum of the capacities constituents. Why the capacity of the compound should be greater, and wh caloric should be more forcibly attracted by an atom of oxygen and an atom rise when united than when separate, I cannot explain. This and other an rine when united than when separate, I cannot explain. This and other an mysteries are no doubt connected with those of electricity, gaivanism, and magnetism.





706. The adjoining figure represents an apfor exhibiting, without danger to the specials detonation of chlorous acid.

707. Into a tube of nearly 8ths of an inch-meter, and sealed at one end, about as smech c of potash is introduced as will rue above the about one inch. The mass thus situated is fused by means of a spirit lamp, or chauffer.

708. The tube, being then charged with

proportion of sulphuric acid, is corked gent suspended within a stout glass cylinder, as drawing. It is then surrounded, near the bott another tube, supplied with boiling water. the hot water is applied only to that part of the which contains the salt; but as soon as the tube is pervaded by a greenish-yellow colo monstrating the evolution of the chlorous at outer tube containing the water is to be rai that the gas may be generally heated by i explosion soon follows, from the influence of spectators are protected by the glass sylinder.

Preparation of Chloric Acid.

709. When a solution of potassa (co potassium) or the carbonate of this all invegnated copi with chlorine, crystals precipitate, which consist of the caid in union with potassa. If to a solution of the chlorate of potassium precipitates. From this an aqueous dilute solution of the desired acid may be obtained by filtration.

116. If, in lieu of a solution of potassa, water holding baryta suspended, impregnated with chlorine, a chlorate of baryta may be procured, from lich the baryta may be precipitated by employing, as nearly as possible, tequivalent of sulphuric acid. It was by this process that Chenevix inversed chloric acid; but it is alleged that when thus procured, it retains

mirate proportion of sulphuric acid.

The Properties.—Chloric acid, thus procured, is inodorous, colourless, ar and astringent. It does not precipitate solutions of lead, mercury, or liver, which, for a great majority of the compounds of chlorine, are infallies tests. When concentrated by evaporation, at a gentle heat, it is releast to an eleaginous consistent, and acquiring a yellowish tint, also an district that of nitric acid. In this state it ignites paper, and other ormais products, and is capable of converting alcohol into acetic acid. It is incomposed by many substances having an affinity for oxygen; and yet, in thing upon iron or zinc, is said to cause the oxidizement of these metals, set, like nitric acid, at its own expense, but by the decomposition of water, if which the hydrogen is in consequence evolved. Many bodies which do tet otherwise react with it, cause its decomposition when aided by the solutions.

Of Oxychloric or Perchloric Acid.

712. Preparation.—After the chlorous acid has been liberated from chloses of potash, the residue consists partially of oxychlorate of potash, as already stated. (700.) This is mingled with bisulphate of potash formed at the same time, but may be separated by repeated solution and crystallization, as the bisulphate is more soluble.*

712 Oxychloric acid may be obtained from oxychlorate of potash by callition in a retort with its own weight of sulphuric acid, diluted with a weight of water at the temperature of 280°. It is purified by carefully presisting the sulphuric acid which comes over with it, by means of ba-

tytic water, and redistillation.

714. Properties.—Like chloric acid, it is insusceptible of the gaseous form, and, as a liquid, exists only in combination with water, being limpid, colories, and having a lively acid taste. It reddens, but does not subse-

Is evolving oxygen from chlorate of potassa, by means of a porcelain retort chauser of coals, it excited surprise, that while the greater part of the gas could welthined in a glass retort, without softening the glass, there was a portion which ised a higher temperature than that which flint glass is capable of enduring. will contemplating some experiments for the explanation of these phenomena, I inmed that Soubieran had furnished the true explanation of the mystery. He had **Installed that one portion** of the salt, receiving two equivalents of oxygen from **Eather, became converted** into an oxychlorate, of which the decomposition was The district than that of the chlorate. Subsequently this process has been resorted by my young friend and late pupil, Mr. Boyé, and my son, to obtain oxychlorate, The mass which remains after all the oxygen has been expartial decomposition. The man which remains and a consists of a mixture of the can be extricated without softening flint glass, consists of a mixture of the call the calls of potassa. potassium and of oxychlorate of potash. As of all the salts of potassa, in contion is the most sparingly soluble in water at 60°, by solution in this said mile boiling hot, and cooling the solution, the salt precipitates, and may be is by separating this part of the process.

quently bleach an infusion of litmus. It is decomposed ther by the solar rays, sulphurous acid, nor sulphuretted hydrogen. It dissolves zinc and iron with disengagement of hydrogen. It exercises strong affinities, and is the most enduring of the combinations of chlorine with oxygen; which is the more surprising, as it is in general true that in proportion as any one ingredient predominates in a compound, it is the more easily separated in part.

715. By the reaction of sulphovinic acid, with oxychlorate of barytes, Mr. Boyé and my son have procured an ether, which in its explosive energy, is scarcely equalled by the chloride of nitrogen. It is I believe the only ethereal compound which is per se explosive, or which detonates from a

mechanical shock.

SECTION III.

OF BROMINE.

716. This name has been given to a substance analo-

gous to chlorine, from the Greek spanes, fetidity.

717. Bromine was discovered by Balard in 1826, at the salt works of Montpelier in France, in the mother waters of marine salt, in the state of bromide of sodium or magnesium. Since then it has been found in the water of the Dead Sea, and in the greater part of the salt springs of the continent, especially those of Germany. In those of Theodorshalle near Kreuznach, a sufficient quantity has been found, to make it profitable to effect its extraction. Common salt, in its natural state, often contains traces of the bromides of sodium or magnesium.

718. Preparation.—The mother water of marine, or common salt, is impregnated with chlorine, until it acquires a hyacinth-red tinge. The chlorine combines with the hydrogen and magnesium of a bromide of magnesium, which exists in that water. The bromine, thus displaced, mingles with the water, which is to be washed with ether. The resulting ethereal solution of bromine, being treated with potash, a bromide of potassium is produced, which, heated in a retort with diluted sulphuric acid and manganese, yields bromine, as chlorine is obtained from a chlo-

ride by like treatment.

719. Properties.—Bromine is a liquid, but is so volatile, that a single drop is sufficient to fill a flask with its reddish-brown vapour. The specific gravity of bromine is 2.966, being nearly three times the weight of its bulk of water. It freezes at a temperature of from 7° to 12° below zero.

It has, when frozen, a crystalline and leafy texture, with a lad-gray colour, and a lustre almost metallic. It boils at the temperature nearly of 89°, forming a vapour resembling that of nitrous acid, and more than five times as heavy as atmospheric air. It does not conduct electricity. Flame settinguished in the vapour of bromine, acquiring a greenic colour previous to its extinction. Bromine is slightly poluble in water. Its solubility is not sensibly augmented by heat. The solution has an orange colour, and emits red fumes. In alcohol it is more soluble, than in water, and in ether still more so than in alcohol.

720. It acts upon vegetable colouring matter and organic products, like chlorine, in general, decomposing them in consequence of its affinity for hydrogen. Bromine forms with starch, a yellow compound. It corrodes the skin, imparting a yellow tinge, which endures till the skin is removated. In its habitudes with oxygen, hydrogen, sulphur, and phosphorus, and the metals, it has a great analogy with chlorine, but generally its affinities are not so strong. From its reaction with potassium, an intense and almost explosive combustion is said to ensue. When taken intermally, bromine acts as a virulent poison.

721. Bromine is supposed to be one of the active substances in mineral springs, especially in those which contain common salt. By means of nitric acid it may be obtained in the form of a deep brown precipitate, from the mother waters in which it exists, but there is much lost by its solution, and subsequent volatilization during the eva-

poration of the solvent.

Experimental Illustration.

722. Bromine exhibited as a liquid; also in the state of vapour.

COMPOUNDS OF BROMINE WITH OXYGEN AND CHLORINE.

Of Bromic Acid.

Bromine forms but one compound with oxygen, called bromic acid,

which was discovered by Balard.

Preparation.—When sulphuric acid is added to bromate of baryta in water, a sulphate of baryta is precipitated, and bromic acid resolution, which may afterwards be concentrated by evaporation.

Properties.—Bromic acid, thus obtained, is a liquid of the consist-

use ox and promise. Bromie acid in the same promise. Bromie acid in the same promise. Bromie acid in the same promise. Bromie acid but it has a strong taste, acid but it has a strong taste, acid but it will be not gen for their radical, decompose the same power acid produces the same power acid cannot exist.

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Of Chloride Bromine.

727. When a current of chlorine is passed through bromine, a liquid compound is produced of a reddish-yellow colour, but not so deep as that bromine. This liquid is volatile, of an intolerable smell, producing that has an excessively disagreeable taste, and a colour resembling that of e chlorine. Water dissolves this chloride, acquiring the power of bleaching litmus. Bases produce with its ingredients, a bromate, a bromide, and chloride.

SECTION IV.

OF IODINE.

728. Iodine has been found in various sea plants, esquially the common sponge, also in mineral waters in a variety of regions of the earth, remote from each other. exists also in combination with various fossils. From the experiments of my late friend Dr. Steel, of Saratoga, a others, it appears to be an ingredient in some of the mineral waters of that place.

729. Preparation.—Iodine is obtained from the lixivit of kelp, from which carbonate of soda is manufacture. After all the soda has been crystallized, the residuum concentrated, and being heated with sulphuric acid, in retort, the iodine passes over, and condenses in shini

crystals of an intense purple or black colour.

730. Iodine may be precipitated from the mother water of salts, with which it is naturally associated, by a mixtre of eight parts of sulphate of copper, and one of green supplied of iron. From this precipitate iodine may be a tained by intense ignition, in a retort, with an equal quality of dry peroxide of manganese.

731. Properties.—When solid, iodine is of a bluish-blacolour, friable, and almost insoluble. It stains the skin y low. It fuses at 225° and volatilizes at 350°, in a beauti

iodine. 131

red. Its taste is acrid and hot, and continues for a time in the mouth. When taken internally, it acts as son. It is incombustible either in oxygen, or atmoic air; but forms acids severally with oxygen, chlound hydrogen, called iodic, chloriodic and iodohydric*

In its habitudes with the Voltaic pile, it is more o-negative than any other matter, excepting oxygen, ir, chlorine, bromine, and probably fluorine. With e varieties of fecula, starch, sago, arrow root, &c., io-produces an intensely blue colour; so that these subes are reciprocally tests for each other. When moisit vaporizes perceptibly, producing an odour similar t of chlorine, but which yet has a peculiar character. pecific gravity of iodine in the solid state is 4.946.

- The vapour of iodine is alleged to have the highest ic gravity of any known aëriform fluid, being 8.716, arly nine times as heavy as atmospheric air. In cong it is peculiarly prone to crystallize, assuming the of an elongated octoedron, with a rhomboidal base. Fr does not dissolve more than 7000 th of its weight, aring a russet colour, but no taste. When the water salt added to it, especially muriate or nitrate of amait dissolves a larger quantity of iodine. The aqueolution does not give out oxygen in the solar rays, estroy vegetable colours. Iodine has a great analogy lorine and bromine, though more feeble in its affinihan either.
- 3. Soubieran recommends that, in order to apply has a test for iodine, the liquid to be essayed should ndered slightly acid by means of nitric acid. After addition and that of the starch, it will, in the course hour, acquire successively a reddish tint, a brownishable, and finally a black colour; or, in other words, lue by its intensity, becomes equivalent to black. It cen alleged that in this way iodine may be detected iquid of which it forms only the 430000 th part.

4. Another mode is to include the liquid to be tested tottle made air-tight by means of a cork, from which pended a piece of moist paper sprinkled with finely

Hem hydriodic has hitherto been applied to this acid, but Thenard, as well kells it iodohydric acid. The considerations which induced me to make will be given hereafter.

powdered starch. If iodine be present, it tinge the starch. It is allowed by Baup that iodine may be thus discovered, when existing in a liquid, in a proportion no

greater than that of a millionth-

735. Balard recommends that, after boiling the liquid with a small quantity of starch, a solution of chlorine in water be added by means of a tube descending to the bottom. The chlorine, at the line of contact, disengages the iodine from its combinations, and enables it to act upon the starch. I resorted to a similar process, about twenty years ago, using sulphuric acid in the manner in which the chlorine is employed by Balard.

Experimental Illustrations.

736. A glass sphere containing iodine, on being warmed, appears filled with a violet-coloured vapour.

737. To a large glass vessel, containing some boiled starch diffused in water, a small quantity of iodine being added, the fluid becomes intensely blue.





739. Heat nearly to the temperature of chullition about two ounces of concentrated sulphuric acid, in a glass globe like that represented in this figure.

789. It is preferable to have the whole of the globe heated, with due caution, over a large charcoal fire. Then quickly traceferring it to the iron tripod, perviously heated, and furnished with a small bed of hot sand, throw into the acid about half a drachm of iodide of potassium, sometimes called hydriodate of potash. Instantaneously the onvity of the globe will become replete with the splendid violet vavour of iodine, which will soon after condense, on those portions of the glass which are first refrigerated, in crystals, symmetrically arranged, of great beauty and unusual size.

133

COMPOUNDS OF IODINE WITH OXYGEN.

Of Iodic Acid.

740. When iodine is subjected to a current of chlorous acid gas, previously dried by passing over chloride of calcium, the gas is absorbed, and a yellow liquid produced. From this, heat expels all the chlorine of the acid,

while its oxygen, uniting with the iodine, forms iodic acid.

741. Properties.—Iodic acid is an inodorous crystalline solid, much heavier than water, with an acid and astringent taste. It deliquesces in most air, but remains unaltered when the air is dry. In water it is soluble, but is precipitated from it by alcohol, in which it is insoluble. Its aqueous solution first reddens and then whitens litmus. With a great number of malifiable bases it forms salts, which detonate if mingled and ignited with any dry combustible matter. In common with bromic acid, it is decomposed by those acids which have hydrogen for their radical, and by many others which have not their highest proportion of oxygen. It contains one stom of iodine, and five of oxygen.

Of Hyperiodic and Iodous Acid.

742. An acid, containing more oxygen than iodic, has been recently discovered by Magnus, to which the name of hyperiodic has been given. But little has been ascertained respecting its properties. Sementini has asserted that he has discovered two additional compounds of oxygen with iodine, one of which he calls oxide of iodine, the other iodous acid. Their existence, however, requires farther confirmation.

Of the Chlorides of Iodine.

743. According to Thenard, chlorine forms with iodine a protochloride and a perchloride. The former contains one atom of each ingredient, the later consists of five atoms of chlorine and one of iodine. The protochlo-

no is the chloriodic acid of Davy.

Thenard awards the appellation of acid to neither. Chloriodic acid is obtained by subjecting iodine in excess, to the action of chlorine. A liquid is produced of a deep reddish-brown colour, much heavier than water, and having in its mechanical properties a great analogy to bromine. It has an acid taste, and reddens litmus. Water dissolves it without sustaining or causing any decomposition, but abandons it to sulphuric ether. If the abovementioned process be so varied as to have an excess of chlorine, per-chloriodic acid is produced, which is a crystalline and volatile substance of a yellowish-white colour, and emitting an effluxium so irritating as to produce tears and a sense of suffocation.

Of the Bromides of Iodine.

745. Bromine combines with iodine in two proportions. A protobromide is obtained when iodine is subjected in excess to the action of bromine. It is solid, and when warmed affords reddish-brown vapours, which condense into crystals of the same tinge, in shape resembling fern leaves. By the same process, when the proportions are reversed, a perbromide results,

which is liquid. Both of these bromides are soluble in water, and I without reddening litmus. Subjected to the action of the Voltaic pike mine goes to the positive, iodine to the negative pole.

SECTION V.

OF PLUORINE.

last edition of this Compendium it was stated, that a In 7. bearing the name at the head of this section, was infer y chemists; and that I had no doubt as to the existence **by** 1 that statement was written. Bandrimont has succeeded resting and energetic element, by passing fluoride of ium, or preferably by heating a mixture of intimately giea chlori coxide of manganese, with concern or calcium and b at this process does not evolve sulphuric : d. It is to be r of purity, in co rine in a que ce of the simultaneous evoluti fluosilicic : n a small proportion. the ibed as a gas of a yellowish-l r.—Fluor 741. F or a prime mingled with a smell of colour, wi TD 00 i but not with glass. The ob-12.4 sugar. it, so far as they extend, justify t tions 1 e DE ap which previous knowledge and re ies. ferences 1 ing had given its pretensions to a place amor e; and go to a Hass. (627, &c.) (633, &c.) halogen bodies or the basacig 748. As there are no known o sounds of fluorine with any of th ments comprised in the class to wnich it belongs, consistently with th rangement to which I have declared my intention to adhere, so for consideration can be given to it, until the bodies are treated of with w

its most important combinations are formed.

SECTION VI.

OF SULPHUR.

749. Sulphur is a mineral production, well known in a merce under the name of brimstone. It is sold but rolls and in flowers. It is found pure in the vicinit volcanoes, of which it is a product. In combination metals it is widely disseminated. From some of its tallic sulphides, which are known under the name of phurets, or pyrites, it may be obtained in the pulver form, to which the name of flowers has been given sublimation.

Properties.—Sulphur is yellow, inodorous, and inbecomes electric by friction, and is liable by the h of the hand to be fractured with a slight noise. It ates and burns with a feeble flame at 180°, and it 225°, and by pouring out the liquid portion, after ass is partially congealed, it may be obtained in is. In close vessels at the temperature of 600° it zes, or sublimes, and afterwards condenses in the nown form of flowers as above stated. The flowers the microscope ascertained to be crystalline, and nerally contaminated by a minute portion of sulphucid, which may be removed by repeated washing.

. All the metals, when presented, in thin leaves or r, to the vapour of sulphur without access of air, into combustion with it, forming compounds which een designated as sulphurets; but which, as I have ought to be called sulphides. (686-7.) Combustion isues when the metals in a divided state are heated ulphur. The sulphides, formed with the metals of rths and alkalies, are soluble in water. From the ing solution the sulphur is thrown down by acids. phosphorus, sulphur is susceptible of a slow as well ick combustion. In consequence of the low temperaat which it is capable of becoming converted by comn into sulphurous acid, sulphur may be burned out of wder without causing it to flash. If raised to the rature of 369°, it enters into a more active reaction. products of the combustion of sulphur are sulphurous mingled with a small portion of anhydrous sulphuric

The hue of the flame when the combustion is slow it; in oxygen its flame is of a splendid purple. Beralleges that when sulphur is rubbed on any body, a for instance, which has been previously warmed, the not sufficiently to inflame the sulphur, an extremely blue flame is produced with a peculiar odour. This he conceives to be the effect of the evaporation, unpanied by any combustion; "since a cold body held it is covered with the flowers of sulphur unchanged." reason, however, appears insufficient; since the sum of one portion of the sulphur does not demonstrate another is not oxydized, any more than the design of carbon upon a cold body exposed to a smoky, proves that another portion of carbon, arising from

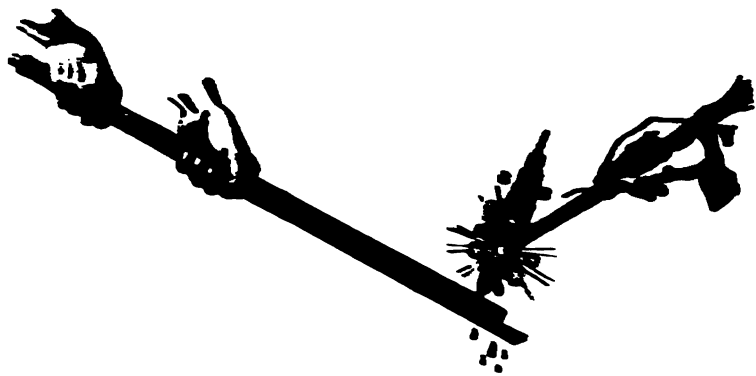
the same source, cannot at the same time be converted into carbonic acid, as is known to be the fact.

752. Some very curious anomalies have been observer suspecting the phenomena of sulphur when kept over the fire after fusion, which the limits prescribed to this would not allow me to introduce.*

Experimental Bustrations.

- 753. Sulphur exhibited in flowers and in rolls; also crys tallized as abovementioned.
- 754. Combustion of Dutch gold leaf and of an iron but by sulphur. Iron wire converted into a sulphide by the vapour of sulphur emitted in a jet from the touch-hale of a gun barrel, made red-hot in the vicinity of the apesture.

The Combustion of Ivan by a Jet of Toporised Sulphes.



733. If a gen-decree he beened red-dut at the decreed and a piece of subplant he there a may it an element the mattrix with a carit, or blowing into it an agment he of vaporised subplant will proceed from the mach-halo Emposed to that a bunch of som were will have as a general in oxygen gotand will full down in the from it theret probables in the same of probabplants. Hydrical if periods, expensed to the sea insest may a subplicit of a few red product.

The latter is designed the different proportions of entry entiring in any animal relativity in the other ingredient. I employ the different members in the different of Themselvania and others.

Organized beig-	Oxilian	Apper Technology	
I man	1 with	PARTED SE	
- 10:00s	1 20:02	siereier yr vieruis.	
E adum	20:ms	esteration is riceral.	

1:	3 atoms	trioxide.
1 atom	4 atoms	quadroxide.
Either 2 atoms	3 atoms ?	- ido
or 1 atom	11/	sesquioxide.

757. The monosyllables di, bi, tri, qua, have an analogous influence on the meaning, when used before any other of the words employed as ove, to distinguish the compounds severally formed by the basacigen elemes, (633,) hence we have dichloride, protochloride, bichloride, trichloa, quadrichloride, &c. It will be perceived that as in the terms quamide and quadrichloride, the monosyllable qua, has such letters added as ry render the resulting epithet easy to pronounce, and agreeable to the ear. 758. The second stage of combination in which the proportion of the etro-negative ingredient exceeds the ratio of equality, has been distininhed by prefixing the word deuto. Hence deutoxide, deutochloride, misbamide, deutiodide. I preser bioxide, as more precise and descripwhere the presence of two atoms is to be indicated. Per is prefixed to gaily the presence of oxygen in a maximum degree, and in the case of m, is used to designate a sesquioxide, in that of mercury a bioxide. But is manosyllable is also prefixed to compounds containing any number of was, whether forming a base or an acid. Hence, we have an acid disspinhed by the appellation perchloric, which contains seven atoms of The syllables in question are prefixed by the French chemists to n words chlorure, bromure, iodure, fluorure, cyanure, as they are by the kinish chemists, prefixed to the modifications of those names which they mploy.

COMPOUNDS OF SULPHUR WITH OXYGEN.

_	With two atoms of oxygen, forms sulphurous acid, equi-			
One atom of	} valent	•	32	
mipher, 16,	walent With three atoms of oxygen, forms sulphuric acid	, equi-		
	(valent	•	40	
_	With two atoms of oxygen, form hyposulphurous	acid,		
Two atoms of	equivalent	•	48	
199 , 32,	equivalent With five atoms of oxygen, forms hyposulphuric	acid,		
	equivalent	•	72	

Of Hyposulphurous Acid.

759. This acid exists only in combination with salifiable bases, and of the sales formed I believe no useful application has been made. Any attend to explain the method in which hyposulphites are produced, will be the salific oxides.

Of Sulphurous Acid.

Preparation.—It is formed by the ordinary combustion of sulphur, by boiling sulphuric acid on sulphur, on mercury, or on any other subtwelly which it may be partially deoxidized.

761. Properties.—Sulphurous acid is a colourless gas, possessing the bown odour of burning sulphur. It is incapable of supporting combining and is deleterious to life, a spasmodic closure of the glottis following any attempt to respire it.

It first reddens and then bleaches litmus, and destroys organic

1.

colours generally. It is used on this account to bleach silk and woolphurous acid is soluble in water, which absorbs 48 times its bulk. It a solution of this gas is exposed to the air, it absorbs oxygen, and is verted into sulphuric acid. This acid, with four times its bulk of a forms a crystalline hydrate, which melts under 40°, disengaging greater part of the acid. After being rendered anhydrous by passing chloride of calcium, sulphurous acid gas, by exposure to a temperatu—12°, condenses into a colourless, transparent liquid, having the sp gravity of 1.45. When dropped in vacuo on the bulb of a spirit mometer, previously at 50°, and surrounded with cotton, the intense of — 90° will be indicated. It is even said that alcohol has been f in this manner. Sulphurous acid gas is decomposed at a red heat, by hydrogen or carbon. It is displaced from its combinations, by a acids except cyanhydric (prussic) and carbonic acid.

Ingregnation of Water with Sulphurous Acid, by means of an appropriate Appr



763. Into the open seed tall receiver, a recurred ; fastened, so as to describe below the me other end of the pipe t in a brass soutet, into wh inserted the stem of an ingless funnel. The placed over the sh pneamatic cistern, cove an inch deep with w includes a stand sup tumbler of the same b pipe, extending from a se pump, rises within the rec nearly as high as the stan under these circumstance pump be put into action equent exhaustion of th from the receiver cases into it of the water from th tern, until the recistance this water opposes to a fi

elevation is greater than opposed by the water is tumbler, to the entrance of air from the recurved pipe communicating will funnel. The air of the funnel will then be drawn into the receiver throug liquid in the tumbler; and if sulphur, carbon, phosphorus, a candle, lamp, o inflammable gas be placed, while burning, under the funnel, the fumes may be to pass through the water, which may be coloured by litmus, or may contain ammonia, baryta, or any other desirable agent, which it may be capable of disse or suspending.

Of Hyposulphuric Acid.

764. This acid is obtained by passing sulphurous acid gas the peroxide of manganese suspended in water in a finely divided state. I mass be kept cold, the peroxide is reduced to the state of protoxide, the oxygen forms with the sulphurous acid, hyposulphuric acid, with the protoxide, produces a salt, which, remaining dissolved, manganetised by crystallization. By the addition of sulphide of barium solution of the resulting crystals, the manganeous is precipitated i state of a sulphide, and hyposulphate of baryta is obtained. From this the hyposulphuric acid may be separated by sulphuric acid, and co trated by evaporation in vacuo, till it acquires the specific gravity of I

By heat or farther concentration, it is decomposed into sulphurous and sulphuric acid.

765. Hyposulphuric acid is a colourless, inodorous liquid, which reddens litmus, has an acid taste, and dissolves zinc with disengagement of hydrogen.

Of Sulphuric Acid.

- 766. Sulphuric acid has been known since the close of the fifteenth century, when it was obtained by Basil Valentine by the distillation of green vitriol, or sulphate of iron.
- 767. Preparation.—This acid may be obtained by burning sulphur and nitre in chambers lined with lead, or by the process abovementioned, by which it was originally obtained; whence the almost obsolete name, oil of vitriol. It is best purified by distillation.

768. I shall defer for the present the illustration of the process for procuring sulphuric acid by sulphur and nitre; also any exemplification of its habitudes with other bodies.

769. Properties.—It is a liquid, oleaginous in its consistency, caustic when concentrated, intensely acid when dilute. When three parts are added to one of water, a boiling heat is produced. (350.) Hot water explodes with it as with a melted metal. It is diluted by the absorption of moisture when exposed to the air. No acid equals it in the power of reddening litmus. When pure it is colourless and has but little smell.

Of the Sulphuric Acid of Nordhausen, and of Anhydrous Sulphuric Acid.

The sulphuric acid of Nordhausen differs from that in use in this country, in containing a portion of acid, which being free from water, is called athydrous. This anhydrous portion being volatile, assumes the form of vapour, and, meeting with the moisture of the air, condenses into the times. (517.)

The furning acid of Nordhausen is obtained by calcination and distinct from sulphate of iron, (known also by the name of green vitriol) contained in retorts of stone-ware. It may be obtained also from white remover sulphate of zinc by similar treatment. The anhydrous acid may be parated from the other portion by gentle distillation, with the aid of a remove diver, previously well desiccated. It is a crystalline body removing asterstos, and may be rubbed between the fingers like wax, when their being attacked. In the air it emits thick fumes having an acid mell. At a temperature above 64° it is liquid. Once congealed it cannot be fused without great care; as the temperature at which it is vaporized, is but little above that at which it liquefies. Hence it is apt to taken a sudden enlargement of bulk which causes it to be thrown out of the cantaining vessel. When vaporized it forms a colourless gas. Neither

in this state par in its crystalline from has it any effect on litmus | sandered perfectly dry. When passed through a red-hot tube of pore it is smolved into oxygen and sulphurous acid.

772. Either canadic lime or baryta enters into a species of comba

with this gas, forming with it a sulphate.

773. The solid anhydrous acid, thrown into water, produces a contion resembling the effect of a hot iron, and, when mingled with an equality proportion of water, explains with a face sufficient to fracture a vessel.

774. The funing acid of Nordhausan is of use for the solution of it employed in dyeing; as the anhydrous acid answers better for this put

then the aqueous.

775. It combines chemically in four proportions with water. The pound committing water in the least proportion is formed in some a processes for producing the acid of Nordhausen. It is a crystalline I which probably consists of 2 atoms of anhydrous acid and 1 of a Pour parts of the anhydrous acid and about 1 of water form the contrast acid of the shops, of the specific gravity of 1.85, which is consist to contain 1 atom of water to 1 of acid. When the acid is to the water to 2, a compound results, of which the density is greater than the 1 density of the constituents, and which probably consists of 2 atom water to 1 of acid. A similar alteration of the density follows the add of water until the specific gravity is reduced to 1.632.

Of the Chlorides of Sulpher.

176. According to Themard, there are two chlorides which are liquids. One contains 2 atoms of sulphur to 1 of chlorine; the other atom of each ingredient. The protochloride is a yellow, vincid, oleaging liquid, heavier than water, and which boils at 250°. The other is rediction, the volatile, furning and acrid, and boils at 147°. Both are deciposed by water and alcohol.

Of the Bromide and Indide of Sulphur.

olenginous, faming liquid. When sodies is bested gently with sulphus torms a brilliant crystalline iodide, of a speci-gray colour.

SECTION VII.

OF SELENIUM.

179. In 1917. Berzelius, examining, in concert with Gahn, the method of preparing sulphuric acid, as practised at Gripsholm, in Swei discovered a sediment in the acid, partly red, partly brown, which, tre by the blowpipe, produced the odour of a rotten radish, and left a mit portion of lead. The odour thus evolved had been considered by Klap as an indication of tellurium. In consequence, Berzeitus took care to lect all the deposition, produced in the manufacture of sulphuric aciding some months; no other sulphur than that of Fahlun being emplo

resulted, to which he gave the name of minn, from the Greek word France, the moon, suggested by its analogy

with tillerium, named from tellus, the earth.

778. Selenium seems much distributed throughout nature. In Sweden it m found combined some imes with copper and silver, sometimes with er only. A small quantity has been detected in cubic galena. In Horway it has been discovered united with tellurium and bismuth; in the hats, combined with lead, copper, and mercury. Stra yer has found it ned with in a mineral from the Lipari is COI

788. Preparation.—From n in which ne actricated by a regia, pro in aq inhurcited hydrogen, re-solution, filtration and evaporation tation by | t, pr on of t reing mass, and sublimation 1 1 1 addition of , penduced by the reaction nium with oxygen in t t wards deoxidized by t ga, is catumated by the potas ang 1 employed, the selenium of the ammonia in the st

blind by the heat.

766. Properties.—Selenium, on cooling after distillation, assumes a hising surface of a deep reddish-brown colour, with a metallic brilliancy tundling that of the blood-stone (hamatite). Its fracture is conchoidal, tons, of a lead-gray, with metallic lustre. Very slowly refrigerated the fusion, its surface become granulated and uneven, of a reddish-gray, and devoid of lustre. By quick refrigeration the characters above indicated mult. Schmirm has little tend :y to crystallize, yet it is capable of separaing in a crystalline pellicle, or of forming a crystalline vegetation, upon the sides of the vessel, from ution in the state of a selenhydrate. When percipitated cold from a cutted solution, whether by zinc or sulphuretted hydrogen, it is red like cinnabar. But if this precipitate be boiled, turn black and consolidates, becoming heavier. When pulverized, seknium becomes of a deep red, and likewise when in very thin layers. With leat it softens, and at the boiling point of water, acquires a semifluidin, becoming completely fluid at a temperature somewhat higher. owing it remains soft for a long time, and may, like heated sealing-wax, the out into filaments. These, by reflected light, are gray, with some metalic brilliancy, but, by transmitted light, are transparent, and of a rubyred colour.

792. When selenium is heated nearly to redness in a distillatory appait assumes, with ebullition, the form of a vapour of a yellow colour, deeper than the hue of chlorine, yet lighter than that of sulphur. report condenses in the neck of a retort in black drops, which coalesce like which are formed by the condensation of mercury. When condensed malogous to the flowers of sulphur, but of a cinnabar-red colour. be mell of a radish is only perceived when the heat is sufficient to be Prictive of oxidation. The specific gravity of selenium is from 4.3, to

Compounds of Selenium with Oxygen.

Scienium has but a feeble affinity for oxygen, yet forms a volatile ide which ! il either of radish or decayed radish. It forms ous and selenic. The latter of these is isomorthe **a.** (474.) K

784. Selenious acid is procured by the co 1 of selenium i

gen gas, or by reaction with nitric, or nitromuratic acid.

785. Selenic acid is obtained by the deflagration of nitre with selin a hot crucible; a seleniate of potash results, which is decomposed intrate of lead, and the resulting seleniate of lead is decomposed to phuretted hydrogen. The sulphuret of lead precipitates, while selenite dissolved in the water employed. When heated to 280 degrees, it is highest concentration, and at 290 degrees is decomposed into a and selenious acid.

786. The highest specific gravity of selenic acid is 2.6. It rest sulphuric acid in its consistency, in its evolving heat by dilution with and in the power of dissolving iron and zinc, with the evolution of I gen. It cannot be rendered anhydrous. When its density is at the With the aid of heat, it or mum, it contains 16 per cent. of v T. t not platinum. With chloro and dissolves copper and even g acid, it constitutes a sort of aqua a, which dissolves both gold ar tinum. Its salts cannot be ed from the sulphates, unless property of detonating with u at a red heat, and that of caus evolution of chlorine, when bo a with muriatic acid. Selenic acid be separated from sulphuric : by aturation with potash, and ig : acia is decomposed into selenium with sal ammoniac. hydrogen of the ammonia.

787. Selenium combines with chlorine and bromine, and with selp

every proportion.

788. As there is not one of the metals which have decided pretant the metallic character, which is not an excellent conductor of both he electricity, and as metallic brilliancy is another striking attribute metallic genus, I cannot understand wherefore selenium, which is admit be destitute of the two first mentioned characteristics, and to possess to imperfectly, should be received into the class of metals; while consider the form of plumbago, is endowed with them all, is excluded cannot consider selenium as a metal. It is stated to have the brilliant hæmatite, which is, I conceive, inferior in that respect to plumbago, the Berzelius considers as pure carbon.

SECTION VIII.

OF TELLURIUM.

789. A metal has been found in the veins of auriferous silver i mines of Transylvania, which has been called tellurium. It is foun in small quantities in Norway, united to selenium. Tellurium has his been discovered in Connecticut. It is found chiefly in the state of an with gold and silver.

with gold and silver.

790. Tellurium displays a metallic brilliancy, and is of a colour be that of tin and antimony, and of a lamellated structure. When me a glass vessel, replete with hydrogen, and slowly cooled, it assumes t pearance of burnished silver. Fused in a vessel, it presents crystal determinable form. It fuses below a red-heat, and above that temps is volatilized. When heated before the blowpipe, it takes fire, and with a blue flame bordering on green, and is dissipated in gray p

nes, which her s the smell of horse-radish. This smell is asited by Berzelius to the presence of selenium. Latterly the same author remes its specific gravity to be 6.2324.

791. Tellurium may be exidized either by combustion or by nitric acid. he mide, exposed before the blowpipe upon charcoal, is decomposed with

ittive violence.

782. Bernelius alleges that tellurium will dissolve in concentrated sulmic acid without being oxidized, in which it differs from other metals. miss that it forms a soluble oxysulphide. The colour of the resulting htien is purplish-red. Tellurium is more especially entitled to our nowen account of its great anale to sulphur and selenium, and of its ming both acids and bases, wh uniting, form telluri-salts. It is upon is ground that Berzelius include it in his amphigen class, and that I conmently place it among the base bodies-

OF RADICALS.

793. Radicals are bodies capable of forming with a baneigen body either an acid or a base, and are divided into those which are metallic, and those which are non-metallic. (633.)

OP NON-METALLIC RADICALS.

The bodies which I place under this head are:—

Hydrogen, Nitrogen, Phosphorus, Carbon,

Boron, Silicon, Zirconion.

SECTION I.

OF HYDROGEN.

794. In its gaseous state, it is the principal constituent al ordinary flame. It is an ingredient in water, and whined with oxygen and carbon, it is found in all vemale and animal substances. It derives its name from

water, and yingui, to produce.

Preparation.—Per sc, hydrogen exists only in the state. In this form it may be obtained by the rediluted sulphuric or muriatic acid with zinc or er of ste a th iron turnings, made red-hot in a gun Lit m: / be evolved in a state of purity, and consequently destitute of odour, from pure water, by Voltaic agency, or by reaction with an amalgam of potassium.

Self-regulating Reservoir for Hydrogen and other Gases.

796. The following figure represents a self-regulating reservoir for hydrogen gas.



797. This very perspicuous engraving can require but litthe explanation. Suppose the glass jar without to costain diglass jar without to com luted sulphuric acid; the inverted bell, within the jur, to contain some zinc, supported on a tray of copper, suspended by wires of the same metal from the neck of the ball. The cock being open when the bell is lowered into the position in which it is represented, the atmospheric air will essape, and the acid, entering the cavity of the bell, will, by its reaction with the zinc, cause hydrogen gas to be copiously evolved. As soon as the cock in closed, the hydrogen expels the acid from the cavity of the bell;

and, consequently, its reaction with the zinc is prevented, until another portion of the gas be withdrawn. As soon as this is done, the acid re-enters the cavity of the bell, and the evolution of hydrogen is renewed and continued, until again arrested, as in the first instance, by preventing the escape of the gas, and consequently causing it to displace the acid from the interior of the bell, within which the zinc is suspended.*

798. By means of apparatus of this kind, I have been enabled to have self-regulating reservoirs of nitric oxide, of sulphydric acid, of carbonic acid, of chlorine, and of chlorohydric acid, merely by changing the materials, and making such a modification of the means of supporting them as the agents employed or evolved require.

The principle of this apparatus is unalogous to that which was contrived by Gay-Lussec. I had employed the same principle, however, when at Williamsburg, Gay-Lussec. I had employed the same principle, however, when at Williamsburg, to moderate the evolution of carbonic acid, before I had read of Gay-Lussec's apparatus. I prefer the modification above described. In the first place, it is internally more easy of access for the purpose of cleansing; secondly, it is much better qualified for containing sulphuret of iron, or marble, for generating sulphuretted hydrogen, or carbonic acid gas; and thirdly, by raising the bell glass, until the liquid within and without is on a level, the pressure may be removed.

In the other form, the pressure on the gas is so great, that, unless the tube, the cock, and the junctures be perfectly tight, there must be a considerable loss of mainrials; since the escape of gas inevitably causes their consumption, by permitting the acid to reach the sinc, or other material employed.

acid to reach the zinc, or other material employed.

Large Self-regulating Reservoir for Hydrogen.



799. This figure represents a self-regulating reservoir for hydrogen, constructed like that described in the preceding article; excepting that it is about fifty times larger, and is made of lead, instead of glass. This reservoir may be used in all experiments requiring a copious supply of hydrogen. When gas is to be supplied to the hydro-oxygen, or compound blowpipe, the perforated knob at the end of the pipe, which has an orifice on one side, is placed under the gallows, G, (seen in the fig. of the compound blowpipe, 331) and fastened airtight to the pipe of that instrument, by the pressure of the screw of the gallows. The gas is retained, or allowed to flow through the pipe, by means of the valve cock, V, which is much less liable to leak, than one of the common form.

Properties of Hydrogen.—It is the lightest of all able substances. One hundred cubic inches weight 3 grains. Its weight to that of oxygen is as 1 to specific gravity, the gravity of air being assumed 0.0689. It is about 200,000 times lighter than platinum. In its y state it smells unpleasantly. When pure it is odour. In its nascent state, as when liberated of an acid, it is extremely prone to take up a portion of sulphur, phosphorus, arsenic, or of some setals. Of the last mentioned property, a most use-ication is now made, which I shall mention when of the process for detecting arsenic.

The respiration of hydrogen, mixed with the same ion of oxygen as exists in atmospheric air, is not by any oppressive sensations; yet a profound mid to have been induced in animals surrounded

by such a mixture. When breathed either in this way unmixed, it will be found to produce a ludicrous alterat in a man's voice, making it shrill and puerile, and so of character as not to be recognised. Sound is said move in this gas with a velocity three times as gr as in the atmosphere. According to the experiments Leslie, the sound of a clock bell was as feeble in hydro By no degree as in air rarefied one hundred times. pressure which has been tried, can hydrogen be conden into a liquid. In consequence of its levity, it esta rapidly from an open vessel, unless inverted. It is I eminently inflammable, yet a taper when immersed in i extinguished. A jet of it, ignited, appears like a fee luminous candle flame, and, if surrounded by a glass to produces a remarkable sound.

802. It has been stated that for equal volumes, all gas have the same capacity for heat; it follows, that for equipments, the capacities must be inversely as their specific gravities or their densities. Hence hydrogen having lowest specific gravity, will have the highest specific has (257.) It is in fact calculated to be as to that of an equipment of air as 13.08 is to 1, and to that of an equipment of water as 3.88 to 1. Its refracting power is and a half times greater than that of the atmosphere.

803. When mixed with oxygen or atmospheric air, subjected to flame, an electric spark, or a wire ignited galvanism, it explodes. With chlorine it explodes un like circumstances, and likewise in the solar rays. burning, it disengages sufficient heat to melt 315 times weight of ice. Dobereiner discovered that platin sponge, a cold metallic congeries, becomes ignited on tering a mixture of hydrogen with oxygen gas, and cause it to inflame by an agency which has not been satisfied rily elucidated. It has since been discovered that particularly elucidated. It has since been discovered that particularly elucidated. It has some been discovered that particularly elucidated in the solar rays.

Experimental Illustrations of the Properties of Hydrog 804. Levity of the gas demonstrated by the ascent of a balloon, or by the effect of filling with hydrogen

glass globe balanced upon a scale beam. (71, &c.) upon the voice shown. Inflammation of a gaseous mixture of hydrogen with atmospheric air by platinated asbestos, or platinum sponge. Apparatus for lighting a candle by a jet of hydrogen from a self-regulating reservoir, either by the electric spark or platinum. (327.) A mixture of hydrogen and oxygen, ignited within a small cannon, ex-

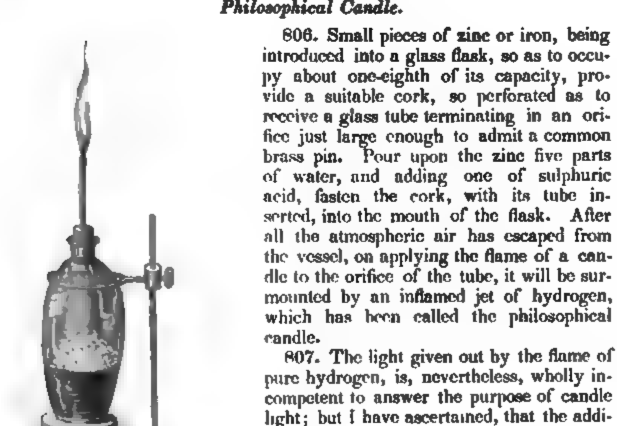


Candle extinguished and re-lighted by Hydrogen.

805. If a lighted candle be introduced into a widemouthed inverted phial, filled with hydrogen gas, the flame of the candle will be extinguished from the want of oxygen. Meanwhile, at the mouth of the bottle, where there is a sufficient access of air, the gas will have taken fire, and will burn with a lambent flame scarcely visible in daylight. Hence if the candle be slowly withdrawn, it will be re-lighted as it passes through the flame.

> tion of a small quantity of spirit of turpentine to the materials obviates this defect.

Philosophical Candle.



Application of Hydrogen and Oxygen in Eur

808. The explosive union of hydrogen with been much resorted to in the analysis of gaseo containing either. For this purpose a stout tub one end, at the other shaped like a trumper drilled into it, near the sealed end, for the introduced into it, near the sealed end, for the introduced other within the tube, for the passage of spark. A known volume of the explosive minimum introduced into the tube, and ignited by a spatelectrophorus or an electrical machine, and the being transferred to a graduated tube, the deby the process is ascertained.

809. The glass tube, employed in this experits appurtenances, is called a endiometer. This was at first applied to the instruments used it sis of atmospheric air, of which one-fifth part gas; but it has since been applied to all instruployed in measuring the results of pneuma analysis. I subjoin an engraving descriptive of the celebrated Volta.

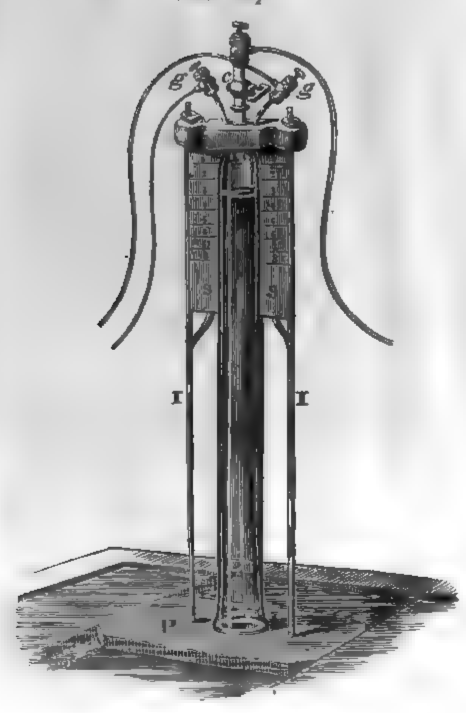
Volta's Eudiometer

510 The endiometer represented by this figure, see next pag by Volta, for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and compounds contained for the analysis of giseous mixtures and contained for the analysis of g

511. The body of this instrument, A. is a cylinder of glass, who had into a brass socket, united by a screw with the cosm. B. This is hollow brass pedestal. C. with the civity in which to be bore of the extes. The glass exlander is also comented into a condition with a cock. By subporting the basis, $\mathbf{F} = \mathbf{T}(x)$ racty of the cosmologic the bore of the cook when open, with this of the lynd of the s the bottom of the basin, the scaled thee, G. graft, by and the w serow out upon a socket, but switch to the consequence. Only hunder, there is a metric obside, there is a metric obside to exceed the consequence. bore of the evin fer environment to the lights of the first light passing the electric some time is now a lower where we for passing the electric some time in laced into the community of a ted this the evince was a great Interpreted to the second Since the William Control Service without it we will will be a Compatible of the conentronity rains to the letter of the

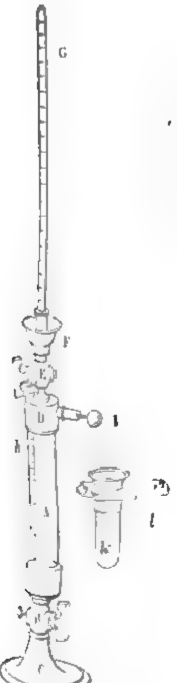
the fire water or the second s

Volumencope.



(Page 149)

catera, the stifes open for the air to escape. Then invert it, the critice being kept



under the surface of the water. Next fill it with the mixture to be analyzed, as for instance a mixture of equal volumes of hydrogen and atmospheric air. Shut the orifice by moving the slide, allow any excess of air to escape, and then, placing the orifice of the measure under the pedestal of the endiometer, open the orifice. the gascous mixture will mount into the cavity of the cylinder. Shut the lower cock, and pass an electric spark through the included mixture. An explosion will ensue, and consequently a portion of the mixture will be condensed into water. By opening the cock, B, the deficit, thus produced, will be compensated by the entrance of an equivalent bulk of water. Open the upper cock, and allow the residual gas to mount into the graduated tube. Detach this tube from the eudiometer, and closing the orifice with the finger under water, before lifting it from the basin, sink it in water, until this liquid be as high without as within the tube. It may now be seen how far the residual air falls short of the 100 measures introduced.

814. It must be evident that we might operate on double the quantity of gas, by taking the measure full of it twice instead of once; and that a mixture of two volumes of air and one volume of hydrogen might be analyzed, by taking three measures equivalent to 300 parts. The loss by the explosion would be the number of degrees that the residue would fall short of 300, when in the graduated tube.

>15 A mixture of three volumes of hydrogen with one of impure oxygen might be analyzed by taking the measure twice full, which is the same as 200 parts. In this case, one-third of the deficit would be the quantity of pure oxygen in 1 of 200, or 50 parts, of the impure gas.

c16 The metallic scale accompanying the cylinder I have never used. Since one of its divisions is equivalent to ten of those on the tube, observations made by means of the latter must be ten times more accurate.

r17. Instead of resorting to an electric spark to produce the inflammation of the gases, I have

the total endrometer a galrano equition apparatus, (335,) by means of which a fewer acture may at any time be agained with certainty.

Of the Volumescope

The lower cond. and is there about this of an inch. There is an obvious at the lower cond. and is there about the lower cond. The tube at the lower cond. The tube at the lower cond. The tube at the consists of a very stout glass tube. At 30 inches in height, and taper. In Camster inside from 2 and 1th to 1 and 1th inches. The least thickness of fine is at the lower cond. and is there about this of an inch. There is an obvious at the choses towards the top, within the space of about 6 inches. The tube take tetween the iron rods 11, which are riveted, at their lower ends, to a simple of the same metal, let into the lower surface of a square piece of plank, because of plank supports the tube, so as to be concentric with an aperture realing with the bore of the tube, and constituting effectively its lower original with the bore of the tube, and constituting effectively its lower original to the same diameter as the end of the tube. Into a perforation in the

centre of the mahogany block, communicating with the bore of the tube, a cook, e, furnished with a gallows screw, is inserted. Through the block, on each side of the perforation, wires are introduced, so as to be air-tight. To the outer ends of these wires two gallows screws, g g, are soldered; to the inner ends a platinum wire, so

as to form a galvano-ignition apparatus. (335.)

819. The apparatus being thus constructed, let it be firmly fixed over the pneumatic cistern, so that the water may rise about an inch above the lower extremity of the tube. To the gallows screws, g g, attach two leaden rods, severally proceeding from the poles of a calorimotor. By means of a leaden pipe, produce a communication between the bore of the cock and an air pump, so that by pumping the air from the cavity of the tube, the water of the cistern may be made to rise into the space thus exhausted of air. On each side of the tube, and between it and each iron rod, there are two strips of wood S S, scored so as to graduate about seven inches of the tube into eight parts. The various distances between these graduations were ascertained by introducing into the tube, previously filled with water, exactly the same bulk of air eight times, and marking the height of the water after each addition. By these means the instrument is graduated into eight parts of equal capacity; and we are by aid of it enabled to measure the gases, and to notice the diminution of volume resulting from their spontaneous reaction, or that which may be induced by the ignition of the wire.

820. The volumescope being so far prepared, and the tube exhausted of air so as to become full of water, close the cock leading to the air pump, and introduce two volumes of pure hydrogen and one volume of pure oxygen, which may be most conveniently and accurately effected by the sliding-rod gas measure. The plates of the calorimotor being in the next place excited by the acid, the ignition of the platinum wire ensues, and causes the hydrogen and oxygen to explode. When they are pure, the subsequent condensation is so complete, that the water will produce a concussion as it rises forcibly against the gum elastic disk, which, aided by the mahogany

block, closes the upper orifice of the tube.

821. If the preceding experiment be repeated with an excess of either gas, it will be found that a quantity, equal to the excess, will remain after the explosion. This is very evident when the excess is just equal to one volume, because, in that case, just one volume will remain uncondensed. By these means, a satisfactory illustration is afforded of the simple and invariable ratio in which the gaseous elements of water units, when mixed and inflamed; which is a fact of great importance to the atomic theory, and to the interesting theory of volumes which hereafter I shall have escasion to notice.

822. Since the accompanying engraving was made, a plate of brass, about a half an inch in thickness, has been substituted for the mahogany block. This plate was made true by means of the slide lathe, the holes for the cocks entering upon the side, and extending inwards and downwards, so as to open into the bore of the tube,

when the plate is in its place.

823. It has been found to contribute much to convenience in manipulating with this instrument, to have a vessel, an iron mercury bottle, for instance, such as represented in page 69, (398,) interposed between the air pump employed, and the volumescope, so as to be exhausted before performing an experiment. Thus assisted, in order to cause the tube to be filled with water, it is only necessary to turn the key of the proper cock. Moreover, by this expedient, the water is prevented from reaching the pump, and when corrosive vapours are produced, lessens the danger of their injuring the mechanism of that instrument.

COMPOUNDS OF HYDROGEN WITH OXYGEN.

Of Water.

824. This liquid may be produced by the combustion of hydrogen gas with oxygen gas. It may be decomposed by passing it in steam over iron, ignited in a gun barrel; also by the aid of acids, by the alkaline metals, by suphurets and phosphurets, by electricity, by galvanism, and by vegetable leaves.

825. Water is necessary to some crystals and to gal-

unic processes. Its powers as a solvent are peculiarly extensive, and are increased by heat and pressure.

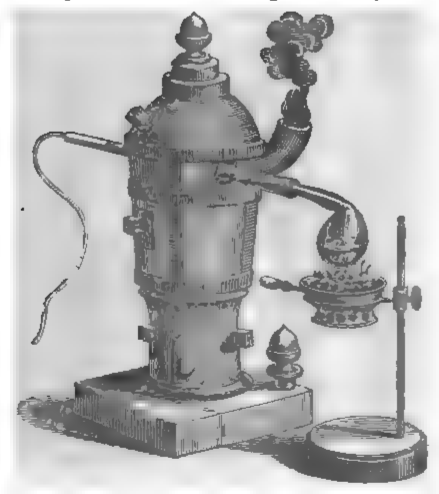
826. Water is one, among other substances, which acts se a acid with powerful bases, while with powerful acids ucts as a base. Berzelius, in some instances, calls it horic acid. It will be seen, as we proceed, that it comhes with various metallic oxides, especially those which constitute the alkalies and alkaline earths. With the lattrespecially it produces much heat in combining, as excoplified in the slaking of lime; and in several of its commations with them, its affinity is too energetic to be wercome by any degree of heat. Excepting acids, any compound in which water exists as an essential constituent, is called a hydrate. Thus slaked lime is a hydrate of lime; but this term is inappropriate, when applied to the compounds which it forms with acids. To them the term quous is applied by Berzelius. The absence of water in my substance in which it is liable to be present, is signised by the word anhydrous. I infer then that its presence should be indicated by means of the adjective hydrous. The vaporization and evaporation of water has, I trust, been sufficiently illustrated. (177, 229, 234.) As a moving power for machinery, as the source of rain, and as the cause of carthquakes, aqueous vapour is, obviously, for good or for evil, one of the most potent agents of nature.

The equivalent of oxygen being And that of hydrogen	
Water is represented by	9

Experimental Illustrations of the Agency of Water.

- 827. No reaction ensues between tartaric acid and carbonated alkali until water is added, when a lively effervescence ensues.
- 828. Concentrated sulphuric acid and zinc remain inactive until water is added, when a copious evolution of hydrogen follows.
- 829. If nitrate of copper be rolled up in tin foil without moisture, the mass will remain inert; but if moistened before it is rolled up, ignition will be produced.

Aqueous Vapour or Steam decomposed by ignited 1



830. Having introduced some turnings of iron or card teeth into an old musket barrel, lute into one it the beak of a half-pint glass retort, about half water; into the other end, a flexible leaden tube. I cover off the furnace, and place the barrel across that the part containing the iron turnings may be e to the greatest heat. Throw into the furnace a t of charcoal and live coals. The barrel will soon! white-hot. In the interim, by means of a chaucoals, the water being heated to ebullition, the st made to pass through the barrel in contact with the iron turnings. Under these circumstances, the oxy the water unites with the iron, and the hydrogen in the gaseous state through the flexible tube.

831. The decomposition of water by sulphurete phurets, and the alkaline metals will be illustrated

time.

Water produced by an inflamed Jet of Hydroge

832. The recomposition of water may be render dent, by means of the philosophical candle, (305,) other inflamed jet of hydrogen, situated within a



Apparatus for the Recomposition of Water.

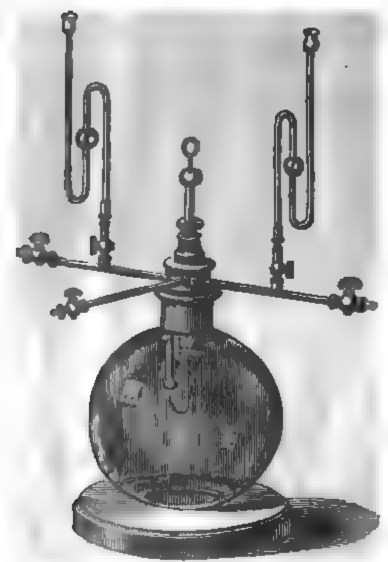


(Page 153.)

iss globe. The glass becomes immediately covered with dew, arising from the condensation of aqueous vapour, odaced by the union of the oxygen of the air with the drogen.

Laurinier's Apparatus for the Recomposition of Water.

8. This apparatus consists of a glass globe, with a neck comented into a brass from which three tubes proceed, severally communicating with an air pump, with reservoirs of oxygen and hydrogen. It has also an insulated wire for proing the inflammation of a jet of hydrogen by means of an electric spark. In w to yet the apparatus into operation, the globe must be exhausted of air, and plied with oxygen to a certain extent. In the next place, hydrogen is allowed to m in a jet, which is to be inflamed by an electric spark. As the oxygen is contain, more is to be admitted. we, was is to be admitted.



If here employed a wire ignited by galvaniam to inflame the hydrogen in this man, and conceive it to be a much less precarious method than that of employ-method machine, or electrophorus. (839).

Develoption of an improved Apparatus for the Recomposition of Water.

This apparatus is represented by the opposite engraving. An inverted bell with a conical neck, is so closed at the apex in the making, as to form a transferoverging cavity, suitable to render the presence of a very small quantity of

ited liquid perceptible to the eye.

I means of the screw rod and plate frame, (248,) this belt glass is secured that position and made air-tight. With the aid of three valve cooks, V V V, the lander pipes, communications with an air pump, a barometer gauge, the sufficiently supplied with oxygen, may be severally opened or closed in. Through a stuffing box which surmounts the plate, a copper pipe, P

in so passed as to occupy the axis of the bell glass, and that of a cil of platings wire, appertaining to a galvano-ignition apparatus, (335, &c.) And copper pipe the minates below in a small platinum tube, and above, outside of the receiver, in a cock C, and gallows screw, by which and a leaden pipe, a communication with a

self-regulating reservoir of hydrogen is at command.

exygen admitted, until the gauge indicates the pressure within the receiver to mearly the same as that of the atmosphere. In the next place, the platinum with being ignited, a jet of hydrogen is admitted, which of course inflames, and continues to burn so long as the supply of the gases is kept up. Soon after the inflamentation of the hydrogen, the resulting water will be seen to coat the interior of the hell glass in drops, resembling a heavy dew, and, continuing to accumulate, will descend in streams into the converging neck of the bell glass. By surrounding this with cold water, the condensation may be expedited, and the deposition of water seen rendered strikingly evident. The gauge employed in this process is that already described. (137, &c.)

- 838. Of the Air in Water. Water naturally contains air. It is to receive the influence of the oxygen of the air thus existing in water, that fishes are furnished with gills, which perform to a certain extent the office of lungs in decarbonizing blood. Fishes cannot live in water which, either by boiling or exhaustion, has been entirely deprived of air.
- 839. The habitudes of other gaseous substances with water will be more advantageously illustrated, when those substances are under consideration.

Experimental Proof of the Presence of Air in Water.

- 840. Water exposed to the action of an air pump, or otherwise subjected to exhaustion, becomes replete with air bubbles.
- 841. Of the Moisture in Air.—Air is not more invariably attendant upon water than water is upon air; nor is the air in water more necessary to fishes, than the water in the air to animals and vegetables. (229, &c.)
- 842. The well known deleterious influence of the winds which blow from the African deserts, arises probably from their aridity. The desiccating power of air is directly as its temperature, and inversely as the quantity of moisture

previously associated with it.

843. There is a certain proportion of moisture, relatively to the temperature, which is most favourable to our comfort. If the moisture be increased without raising the temperature, or the temperature be increased without an accession of moisture, we are incommoded. In the one case, the skin becomes unpleasantly dry; in the other, the air is too much encumbered with aqueous vapour, to allow

perspiration, whether sensible or insensible, to proceed with sufficient freedom.

844. Stove rooms are oppressive on account of the too great aridity of the air in them; and hence the well known remedy of a basin of water, placed upon the stove to furnish moisture by its evaporation.

845. Hygrometric Process of Dalton.—The dew which is observable m vessels containing cold water, in warm weather especially, arises from

the condensation of the aqueous vapour in the air.

=16. According to Mr. Dalton, the less the degree of cold requisite to produce this phenomenon, the greater the quantity of moisture in the air. Hence, by ascertaining the highest temperature at which the water is capabe of producing the condensation, the quantity of moisture may be known]

from a table which he has constructed. (229, &c.)

-47. Daniell's Hygrometer.—Mr. Daniell has contrived an hygrometer upon the principle thus suggested by Dalton. Vaporization is ingeniously assized to produce cold in one bulb of the instrument, in consequence of the cold produced by the evaporation of ether in another bulb, as in the cryopherus. (407, &c.) Two thermometers accompany the instrument, one whin the bulb refrigerated by the vaporization; the other so situated as to indicate the temperature of the atmosphere. As the quantity of aqueous vapour in the air diminishes, the depression of temperature necessary to the precipitation of moisture on the refrigerated bulb increases. The extent of the degreesen is ascertained by the thermometers, the quantity of water in the air liv reference to a table.

-1-. Organic Sensibility of the Beard of the Wild Oat (Avena Sensiture to Moisture.—Hygrometers have been made which are dependent upon the contraction or dilatation which catgut, whalebone, and other subsames of a like nature undergo, in proportion to the quantity of moisture to the air. Among instruments of this kind, that formed by means of the wari of the wild out is pre-eminent for its susceptibility to the influence of treesure. Breathing on it through a minute hole in the case, causes the indications of hygromerce this constructed are not referrible to any standard, agreeably to which a comparison can be made between the dryness of the air in dif-Frent places at the same time, or in the same place at different times.

49. Hygrometric Process by means of a Balance.—It may be presumed that the quantity of moisture in the air is inversely as the weight of

water which will in a given time evaporate from a moist surface. If this presumption be correct, the little square dish here represented may, with the aid of a delicate scale beam, be used as an hygrometer. If it be suspended to the balance, and equipoised while containing a little water, the counter-weight will in a few minutes preponderate, in consequence of the loss by evaporation.

56. The loss of weight within any known period being determined, the reporting power of the air will be as the loss of weight; but as the evaporation is more or less rapid in proportion as there may be more or less agitation, it will not be right to infer that the quantity of aqueous vapour in the atmosphere is inversely as the rate of evaporation, unless the process were uninfluenced by the wind. Of course the dish should be of conve dimensions, accurately determined; 2 inches square for instance.

Compounds of Chlorine with Water.

851. Hydrate of Chlorine.—Berzelius observes that chlorine furnishe the only instance of an elementary substance capable of entering into co bination with water. I allude here to a crystalline compound formed passing the gas through that liquid at a temperature below 40° F. T hydrate thus formed is capable of being sublimed from one part of the containing vessel to another, in consequence of a slight diversity of temperature. It consists of one volume of chlorine, and twenty volumes of aqueous vapour.

852. Solution of Chlorine in Water.—The same eminent author alleges that, in order to obtain a saturated solution of chlorine in water, it is necessary, in the first instance, to expel from the latter all the atmospheric

air.

Of the Deutoxide or Bioxide of Hydrogen, or Oxygenated Water.

853. In 1818, Thenard discovered that water might be made to receive an additional quantity of oxygen, by dissolving deutoxide of barium in liquid muriatic acid, precipitating the baryta by sulphuric acid, and the chlorine by silver.

854. Properties.—The bioxide of hydrogen is as liquid, and as devoid of colour as water. It is nearly inodorous, whitens the tongue, inspissates the saliva, and tastes like some metallic solutions. Applied to the skin, it creates a smarting sensation, more durable in some persons than in others. Its specific gravity is 1.452. Hence, when poured into water, it descends through it like syrup, but is dissolved by agitation. As it is less easy to vaporize than water, it may be separated from that liquid, by exposure in vacuo over sulphuric acid. (309.) In its most concentrated form, it has not been congealed by any degree of cold to which it has been subjected. The most surprising property of this substance is that of giving off oxygen explosively, on being brought into contact with substances which do not unite with either of its ingredients. Thus it explodes by contact with finely divided silver, platinum or gold, and still more actively with oxide of silver or peroxide of lead. The difficulty of explaining these phenomena has already been noticed. (421, &c.)
855. When mingled with the mineral acids, its liability

to decomposition is diminished. If exposed to heat in its

nost concentrated state, a few grains create a violent explosion. When, by dilution with 20 parts of water and exposure to heat, it loses all the oxygen which it holds beyond the quantity necessary to the composition of water, as much oxygen is found to be evolved as the hydrogen in the residual water retains. Hence it is generally supposed to consist of one atom of hydrogen and two of oxygen.

Remarks on Nomenclature.

556. Some of the most eminent European chemists have, most errone-casive and inconsistently, designated the acids formed by hydrogen, with the dictronegative, or basacigen bodies, as hydracids; while analogous compands, formed by other radicals, were designated by prefixing syllables adicative of the electro-negative ingredient. Thus we have had hydrochloric, hydrobromic, hydroiodic, hydrofluoric, hydrocyanic, &c., to signify the acid compounds of hydrogen with the halogen elements; while we have had fluoboric and fluosilicic to signify acids formed with the radicals boron and silicon by fluorine. Thus the former series is characterized by letters taken from the radical, the latter by letters taken from the electro-negative or basacigen ingredient, while hydrogen is placed by the side of oxygen, with which, in properties, it is extremely discordant. (633, 636.)

57. This error I pointed out in an article published in the Journal of Pharmacy, in the autumn of 1833, and in a letter to Professor Silliman.

* The following passage is in the letter to which I have referred.

" In common with other eminent chemists, Berzelius has distinguished acids in which expects is the electro-negative principle, as oxacids, and those in which hydiegen is a prominent ingredient, as hydracids. If we look for the word radical in == : : : : contents of his invaluable Treatise, we are referred to p. 218, vol. 1st, *Ler- we find the following definition. "the combustible body contained in an acid, reasonable base, is called the radical of the acid, or of the base." In the second verge 1.3, he defines hydracids to be "those acids which contain an electroto zwise body combined with hydrogen;" and in the next page it is stated, that "hycrace we divided into those which have a simple radical, and those which have a Composed radical. The second only comprises those formed with cyanogen and sulprogramment Again, in the next paragraph, "no radical is known that gives more धारक जार अर्थ with hydrogen, although sulphur and iodine are capable of combining That is exact proportions. If at any future day more numerous degrees of acidibeaten wra sydrogen should be discovered, their denomination might be founded on the same principles as those of oxacids." Consistently with these quotations, all the their -tegative elements forming acids with hydrogen are radicals, and of tours. by he own definition, combustibles; while hydrogen is made to rank with raygen as an activitying principle, and consequently is neither a radical nor a com-We ble. Yet, page 1:29, vol. 2d, in explaining the reaction of fluoboric acid with n which case fluorine unites both with hydrogen and boron, it is mentioned ** Estance among others in which fluorine combines with two combustibles.

Fig. of opinion that the employment of the word hydracid, as co-ordinate with small must tend to convey that erroneous idea, with which, in opposition to his contained, the author seems to have been imbued, that hydrogen in the one ciam, plays the same part as oxygen in the other. But in reality, the former is emi-

seatly a combustable, and of course the radical by his own definition.

Plane of hydracids, but, with greater propriety as I conceive, places them under sames indicating their electro-negative principles. Thus he arranges them as oxygen acids, ciderine acids, bromine acids, iodine acids, fluorine acids, cyanogen acids, uphur acids, scienium acids, and tellurium acids. These appellations might, I

Afterwards I had the satisfaction of observing, that, in an edition of his Traité, then in the press, Thenard was acting upon a similar view of this subject, and employing the language which I had suggested. Moreover I found that Dr. Thomson had not arranged the acids alluded to under the name of hydracids, but had put each of them under the name of its electronegative ingredient. Hence they were treated of under as many heads as there are basacigen bodies. Or, to be more particular, they were treated of as oxygen acids, chlorine acids, bromine acids, iodine acids, fluorine acids, cyanogen acids, sulphur acids, selenium acids, and tellurium acids.

858. Consistently with the process of abbreviation by which oxacid has been employed to designate an acid formed by oxygen, and hydracid to signify an analogous combination formed with hydrogen, I have made the following abbreviations of the appellations employed by Thomson:—

For	Oxygen acids	to use	Oxacids.
22	Chlorine acids	77	Chloracids.
22	Bromine acids	"	Bromacids.
22	Iodine acids	"	Iodacids.
22	Fluorine acids	"	Fluacids.
"	Cyanogen acids	• -	Cyanacids.
22	Sulphur acids	22	Sulphacids.
	Selenium acids,	"	Selenacids.
"	Tellurium acids	"	Telluracids.

859. The acids formed by oxygen received their names, for the most part, before the basacigen bodies were recognised as elements, or the existence of some of them discovered. Hence, in the case of the oxacids, it is neither customary nor expedient, to prefix any syllables indicating their basacigen ingredient. Consequently, we have sulphuric, selenic, telluric, chloric, bromic, iodic, &c. &c. instead of oxysulphuric, oxyselenic, oxytelluric, oxychloric, oxybromic, oxiodic, &c. The syllables were employed prior to the recognition of the elementary character of chlorine, to designate an oxacid with an extra proportion of oxygen. Thus chlorine was

think, be advantageously abbreviated into oxacids, chloracids, bromacids, iodacids, fluacids, cyanacids, sulphacids, selenacids, telluracids.

"I had formed my opinions on this subject before I was aware that Dr. Thomson

had resorted to this classification.

"As respects the acids individually, I conceive that it would be preferable, if the syllable indicating the more electro-negative element had precedency in all, as it has in some cases. The word hydrofluoric does not harmonize with fluoboric, fluosilicic, fluochromic, fluomolybdic, &c. Fluorine being in each compound the electronegative principle, the syllables, indicating its presence, should in each name occupy the same station. These remarks will apply in the case of acids formed with hydrogen by all principles which are more electro-negative. Hence we should use the terms chlorohydric, bromohydric, iodohydric, fluohydric, cyanhydric, instead of hy-

drochloric, hydrobromic, hydriodic, hydrofluoric, hydrocyanic.

"As by the British chemists the objectionable words have not been definitively adopted, the appellations muriatic and prussic being still much employed, it may not be inconvenient to them to introduce those which are recommended by consistency. In accordance with the premises, the acids formed with hydrogen by sulphur, selenium, and tellurium, would be called severally sulphydric, selenhydric, and telluhydric acid. Compounds formed by the union of the acids, thus designated, with the bases severally generated by the same electro-negative principles, would be called sulphydrates, selenhydrates, and telluhydrates, which are the names given to these compounds in the Berzelian nomenclature. Influenced by the analogy, a student would expect the electro-negative ingredient of a sulphydrate to be sulphydric acid, not a sulphide. The terminating syllable of this word, by its associations, can only convey the conception of an electro-positive compound."

micalled oxymuriatic acid, being supposed to be an oxide of an unknown adical, with an extra dose of oxygen. (888.) At this time, oxychloric acid designates the acid which has more oxygen than the chloric acid.

860. The analogy between the acids formed by hydrogen with the halogen hodies, chlorine, bromine, iodine, fluorine, and cyanogen, render it both desirble and practicable to treat of them in a body, mainly by reference to chlorohydric acid. Hence I shall employ the word halohydric, to designet those acid compounds; and in obedience to similar considerations, the tempounds formed with hydrogen by the amphigen bodies, sulphur, sele-

mm, and tellurium, will be designated as amphydric acids.

**Sol. The compound formed by the union of hydrogen with oxygen, the putoxide of hydrogen, (water,) ought not to be included under the head of the amphydric acids. Of this oxide, the pretensions to the characteristics of a base, are at least as high as those which can be advanced for it as an acid. Of course it cannot, with propriety, be classed with any acid compounds. It is in reality an anomalous substance, performing a part in nature of such pre-eminent importance, as to merit to a certain extent an isotated position, and undivided attention.

862. Names of the halohydric acids, or those formed by the five halogen bodies, chlorine, bromine, iodine, fluorine and cyanogen, with hydrogen, as heretofore given by the French chemists, also by Berzelius, Turner, and others, contrasted with those now employed in this Compendium, agreeably to the practice of Thenard, and with the approbation of Berzelius.*

For hydrochloric use chlorohydric.

"hydrobromic "bromohydric.

"hydroiodic "iodohydric.

"hydrofluoric "fluohydric.

"hydrocyanic "cyanohydric.

863. Names of the amphydric acids, or acids formed by the amphygen bodies of Berzelius (excepting oxygen) with hydrogen.

For hydrosulphuric use sulphydric.
,, hydroselenic ,, selenhydric.
,, hydrotelluric ,, telluhydric.

I cheerfully admit that it would be preferable to employ the word chlorohydric, intend of hydrochloric. My motive for retaining this last, was, that I was unwilling a venture upon a new nomenclature in a language foreign to me, in which it was inexpedient to make changes which could be avoided without inconvenience. I also spec with you, that we ought not to use combustible and oxidable, as having the same meaning. I have deserved your strictures for this inconsistency in my language; but I must suggest as an apology, that the two words were formerly used syntappaces, and that the work, in which you have recently noticed this overlight, was first published in 1806, having been from time to time remoulded for new lattices, without its having been possible to eradicate all that has not kept pace with the progress of science.

COMPOUND OF HYDROGEN WITH CHLORINE.

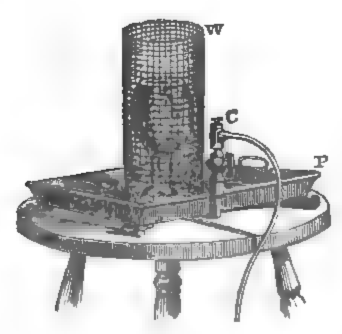
Of Chlorohydric or Muriatic Acid Gas.

volumes they combine spontaneously. In the dark, of where the light is feeble, the union is slowly accomplished but, in the solar rays, takes place explosively. According to Silliman, the direct rays of the sun are not necessary to produce the result. The mixture may also be exploded by the electric spark, or by contact with any ignited matter However the union may be effected, chlorohydric or muriatic acid gas is produced, without any reduction of volume if no water be present.

Synthesis of Chlorohydric Acid Gas.

865. In order to demonstrate the ratio in which chlorine and hydrogen combine, it is only necessary to introduce and ignite in the volumescope over water, equal measures of each gas. If they be pure, there will be a complete condensation. The experiment is conducted precisely as in the case of oxygen and hydrogen, excepting that in lieu of a half volume of oxygen, a volume of chlorine is supplied from a self-regulating reservoir. (798.)

Explosive Reaction of Hydrogen with Chlorine, under the influence of the Solar Rays.



866. A flask is half filled with chlorine over the pneumatic cistern in the usual way, and then transferred to the pan P, so as to have its orifice exactly over that of a pipe which, at the other end, communicates with the cock C, to which is annexed a flexible pipe extending to a self-regulating reservoir of hydrogen. (799.)

567. The flask is surrounded by a wire gaaze, W, and just before the explosion is desired, hydrogen from the reservoir is allowed to occupy that portion of the cavity which was previously unoccupied by the chlorine. It

should be understood that the pan, during this operation, retains a sufficient stratum of water to cover the mouth of the flask, and that this is occupied with the same liquid in part until it is displaced by the hydrogen.

*6*4. The preliminary arrangements being made, a mirror must be placed

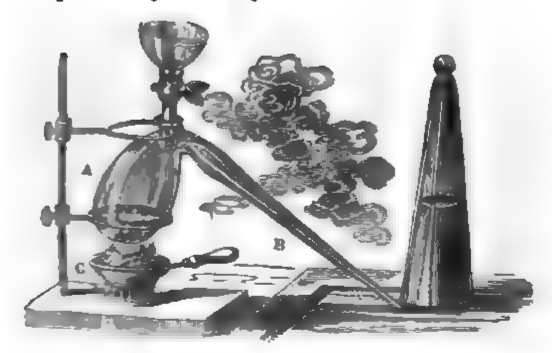
ma situation to receive the solar rays without passing through window glass, and to select them upon the flask. The result is an explosion, from the effects of which the spectators are protected by the wire-gauze.

869. It must be obvious that this experiment can only succeed when the

ster is encloseured.

87%. It should be understood that the condensation arises altogether from the shorption of the gas by the water. (866.)

Preparation of Chlorohydric or Muriatic Acid Gas.



871. Into a tubulated retort, introduce about as much chloride of sodium (common salt) as will occupy nearly one-third of the body, A. Lute a glass funnel, furnished with a cock, into the tubulure. Let the orifice of the beak, Libe so depressed below the surface of the mercury in the cisters, as to be under a bell glass, filled with, and inverted over, the mercury, and properly situated for receiving any gas which may escape through the beak. Prepare about three-fourths as much strong sulphuric acid by weight as there may be salt in the retort. After pouring about onethird of the acid into the retort, close the cock of the fun-≥: the mixture will rise in a foam, and a portion of gascommatter will pass into the bell. As soon as the foam whides, add more of the acid until the whole is introdeced. Then as soon as the foam again subsides, apply the chauffer. C. and chlorohydric acid gas will continue to be copiously evolved. I have of late substituted for the funnel a glass tube of about a half an inch in bore at one end, tapering, at the other end, to an orifice of about the eighth of an inch in bore. This tube, being inserted into the retort through the tubulure, and luted thereto air-tight,

affords a channel for the gradual introduction of the acid which, surrounding the lower orifice of the tube, preven

the gas from escaping.
872. Rationale of the Process.—The water comb with the sulphuric acid is decomposed; its oxygen, with the sodium, forming soda, with which the st acid combines, forming sulphate of soda. The hydright of the water and the chlorine escape as chlorohydright

gas.

873. Properties.—Chlorohydric acid has all the attri-butes of a gas. It is colourless, and, although less active than chlorine gas, is to the organs of respiration intolerably irritating, and if not very dilute, deleterious to life. On escaping into the air, it produces white fumes, from in meeting with moisture. Its affinity for water is so great that this liquid will take up 420 times its bulk, and when in this state, ice is liquefied as if surrounded by fire-When brought into contact with the metals which ecompose water, its hydrogen is liberated, while the chlorine unites with the metal. Equal weights of potassium separate the same weights and volumes of hydrogen from chlorohydric acid, and from water; a result conformable with the inferred atomic composition of both. Presented to metallic oxides, a reciprocal decomposition ensues; the hydrogen unites with the oxyen generating water, the chlorine with the metal producing a chloride. If mingled with oxygen and exposed to the action of heat or a succession of electric sparks, gaseous chlorohydric acid is partially decomposed. This result cannot be extended to more than th of the whole volume. At the temperature of 50°, and under a pressure of forty atmospheres, it becomes a colorless liquid.

874. Its specific gravity is 1.2694, a mean between that of its constituents. The weight of 100 cubic inches is

39.36 grains.

One atom of chlorine, equivalent And one atom of hydrogen, equivalent		
Constitute one atom of chlorohydric acid gas, equivalent	- 37	

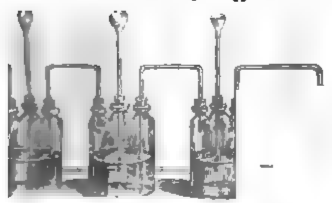
Experimental Illustrations.

Equal volumes of hydrogen and chlorine, being and subjected to the solar rays, (867,) or galvanic, (818,) explode and form chlorohydric acid gas. Gas collected over mercury in tall jars. Water, d by litmus, being introduced, rapidly changes to a sur, and causes the disappearance of the gas. Same reduced by ice, which is rapidly melted.

uration of Liquid Chlorohydric or Muriatic Acid.

It may be obtained by saturating water with the Voulse's apparatus. (See the following figure.) The is nearly pure in all the receptacles excepting the





this figure Woulfe's apparatus is depicted in an improved form. The i in the retort, first passes into the globe where any vapour which may a condenses. It then proceeds along the tube which establishes a commuch the bottle next to the globe. As that mouth of this tube which is bottle, is below the surface of the liquid placed there to absorb it, the gas is up through the liquid, so as to promote its own absorption by the aginideced. It then ruses above the surface of the liquid, where a further takes place. The excess of gas, beyond the quantity absorbed by the lifest bottle, passes, by means of the connecting tube, to the second bottle, rer portion is not there absorbed, reaches the third bottle, in the case of moress proceeds as in that of the first two. Should any of the gas escape wites, it may, by lengthening the last tube, be conducted under a bell with water on the shelf of the hydro-pneumatic cistern, so as not to answer is nearly saturated.

poung the extrication of gas to cease before the liquid in the first bottle, the absorption continuing, the liquid in the second bottle might be transferst, in consequence of the rerefaction of the residual gas rendering it it to resist the atmospheric pressure. In like manner the contents of the might be transferred to the second. To prevent these inconveniences, each bottle a straight tube fastened air-tight into an intermediate neck, ding into the liquid. By these means an adequate pressure is opposed to of the gas, and yet any diminution of pressure, arising from absorption, pensated by the ingress of atmospheric air, ere the liquid can be drawn in next bottle. To prevent absorption from the first bottle into the globe, so, for the introduction of the acid, a trumpet-mouthed tube of small through and luted into the tubulure by a cork with lead and a gum and terminating in a small orifice near the bottom of the retort inside.

888. Of late I have resorted to the following expedient. The beaks of four tuber lated retorts, are drawn out by heating them in a hele opened by a poker in an antiferedite fire, until the beak, by its own weight, is made to extend steelf into a lung tageting take. At the moment when this takes place, by lifting it from the fire and healing the bedy of the retort in a suitable position, the tapering portion of the best benge down, making the desired angle with the other part of the beak. Of course it retains this form when cold. The retorts thus prepared, are so associated that the beak of No. 1, the larger retort, may enter No. 2, through the tabulars of No. 2, and that the beak of this may in like meaner reach into No. 3. Of course a fourth and a fifth entert may, if requisite, he thus made to communicate. The brake are to be lated to the tabularse; and No. 1, being supplied with the salt, and farmaked with a tapering tube for the introduction of the sulphuric said, the process is to be conducted as already described. (673, &c.)

tube for the introduction of the sulphuric acid, the process is to be conducted as all ready described. (873, &c.)

831. Commercial chlorohydric acid is so cheap, that I have found it preferable to use it in the first retort, instead of salt. The addition of sulphuric acid causes the gas to come over pure, without heat at first, but with the aid of a gentle heat, nearly the whole may be evolved, and of course absorbed by the water, placed purposely within retorts, No. 9 and 3. It is preferable to add a fourth retort, and to have No. 3 quite small, holding only a small quantity of water, just adequate to washout of the gas any sulphuric acid which may attend it in the form of a apray—It may be remarked, that one advantage of this process is, that the iron which is unusually an impurity in liquid chlorohydric acid, forms a compound with sulphuric acid, which is not like the chloride of that metal, volatile. Consequently, by thus process, the acid is departed of iron.

862. Liquid chlorohydric acid may also be obtained by distilling a solution of chloride of codiming water with sulphuric acid. In this way there is no used of all apparatus for preferables absorption, as described in the preceding article. The acid course over and collisiones in union with the requisite quantity of water

883. Properties of the Liquid Chlorohydric Acid.—When concentrated, it produces suffocating fumes from the escape of gas. When pure, it is colourless, though usually straw-coloured from the presence of a minute portion of iron.

884. Dr. Thomson informs us that the strongest liquid acid which he could obtain, consisted of one atom of acid, equivalent 37, united with six atoms of water, which being equivalent to 54, the proportion of acid to water by weight was as those numbers, or nearly as 2 to 3.*

* The relative equivalent proportion of chlorobydric acid and water, or prop of said soid, by weight, in aqueous solutions of different specific gravities, i learned from the following table. (See Thomson's Principles of Chemistry.)

Atoms of Acid.	Atoms of Water.	Real Acid in 180 of the Liquid.	Specific Gravity,
1	6	40.659	1.203
t	7	37.000	1.179
1	8	33.945	1.162
1	9	31.346	1.349
1	10	29.134	1.139
1	l n -	27.206	1 1285
1	12	25.517	1.1197
1	18	24.026	1.1127
1	111	14/700	1.1060
1	15	21 512	1 1008
1	16	20.442	T-OHN90
1	17	19.474	L_{00000}
1	18	18.590	1.0860
1	19	17.790	1.0890
1	20	17.061	1.0780

Experimental Illustrations.

quid chlorohydric acid exhibited; also its reacother bodies.

Theory of the Nature of Chlorine and Chlorohydric Acid.

rohydric acid was deemed to be a compound of oxygen with rn radical. When distilled from red oxide of lead, or black ganese, it was supposed to combine with a portion of the oxyoxides, forming oxygenated muriatic acid, the name then rine. To the oxygen thus imagined to exist in it, the activity as a supporter of combustion and as a solvent of metals, was has since been proved that neither carbon nor the metals are n intensely ignited in dry chlorine. The metals are converted, while the carbon undergoes no change. Chloride of sulphur of phosphorus, which result from saturating these substances wine, are devoid of acidity; but the addition of water converts muriatic and phosphorous acid, the other into muriatic and rus acid.

slorine be muriatic acid oxygenated, the discovery of the hypoorous, chloric, and oxychloric acids must establish this anone radical of muriatic acid, by successive additions of the same inciple, gains, loses, and regains acidity, forming first an acid, de, and finally four acids. I have said it forms an oxide, bene must be deemed an oxide, having no acid properties.

see stated, page 156, that Thenard oxygenated the water in tic acid; yet this did not convert it into a solution of chlorine. ceably to the doctrine now universally sanctioned by chemists, acid, consisting of chlorine and hydrogen, is deprived of all those processes by which it was formerly supposed to be

Of Bromohydric Acid.

obtain bromohydric acid, Berzelius recommends that phosphoe placed in contact with bromine under water. The resulting esolved into phosphoric acid and bromohydric acid gas. The ecollected over mercury, or made to produce liquid bromohyunion with water, exactly by the same means as have been ilthe case of chlorohydric acid, which the bromohydric acid much Bromohydric acid is a colourless gas, in smell similar to chloth that a specific gravity of 2.7353. When brought in cone air it produces thick fumes. It is decomposed in passing the heated red-hot. It is composed of one atom of hydrogen fromine.

Of Iodohydric Acid.

ording to Berzelius, in order to procure iodohydric acid, nine ne and one of phosphorus should be placed in contact at the botze or small matrass, and protected from the air by powdered de of phosphorus is formed, which is resolved into phosphoric lohydric acid gas by the gradual affusion of a small quantity of e gas cannot be collected over water or mercury, as it acts on

the one and is absorbed by the other.

then, by means of tubes descending three their postures which is analogous to the mode, already a strated on a large scale, for collecting chlorine. (666.) This process is even more practicable in the case in point; since iodohydric acid is the heaviest gas known, having a specific gravity of 4.3854, or more than four times as great as that of atmospheric sir. In composition and general properties it resembles chlorolydric and bromohydric acid.

892. The compound of hydrogen with flacrine, forming the acid of flac spar or fluckydric acid, improperly called hydroduoric acid, will be defer red for consideration, until boron and silicon have been treated of

COMPOUNDS OF HYDROGEN WITH SULPHUR.

893. It appears probable that hydrogen and suphur may combine in various proportions. Only two compounds, however, have been sufficiently distinguished to be worthy of a place in this work. One of these is a definite compound of hydrogen and sulphur, containing an atom of each, and has hitherto been called sulphuretted hydrogen, especially by the British c emists. The other contains one atom of hydrogen, with a plurality of atoms of sulphur, which, according to Thenard, may extend to the proportions of four, six, or eight atoms to one. To this he has accordingly given the name of polysulphuret of hydrogen.

894. Pursuant to the nomenclature of Berzelina, all the electro-negative compounds of sulphur are called sulphides and are designated by attaching, as an adjective, their radical, with the last syllable changed into ique in French, on ic in English; as, for instance, sulphuretted hydrogen is called by him in French, sulphide hydrique, which in English is rendered by hydric sulphide. This gas has by some chemists, especially the French, been called hydrosulphuric acid, by analogy with hydrochloric acid. The term kydrosulphuric is objectionable from its conveying the idea of aqueous sulphuric acid; hydro being used to imply the presence or influence of water. I have already pointed out the inconsistency of designating some acids by giving precedence to the syllables representing their radical, as in hydrochloric, hydriodic; while in others, the syllable indicative of their electro-negative ingredient has the precedence, as in fluosilicic, fluoboric, chlorocarbonic, and chlorocyanic. If sulphuretted hydrogen is to receive a new name, I would prefer to call it sulphydric acid, as already suggested. (858, &c.)

Of Sulphydric Acid or Sulphuretted Hydrogen.

895. Few persons are unacquainted with the unpleasant odour which results from the washings of a gun barrel, made foul by the explosion of gunpowder, or that produced by putrid eggs. This odour arises from a compound consting of one atom of hydrogen and one atom of sulphur. The celebrated sulphur springs of Virginia are indebted for their odour, and mainly for their efficacy, to this compound; to which the celebrated Thenard has given the name of sulphydric acid.

896. Preparation.—This gas is copiously evolved by the reaction of diluted sulphuric acid with sulphuret of iron. In order to have a supply of it at command, it is only necessary to substitute this last mentioned substance for zinc in the self-regulating apparatus employed for hydrogen,

already described. (706.)

897. As it is absorbed by water and gradually decomposed by mercury, Berzelius recommends that it should be received over brine. Its purity is demonstrated by its complete absorption by a solution of caustic potash, and by its

not rendering lime-water milky.

through water as otherwise it is liable to be contaminated by the generating materials. When the acid is sufficiently diluted, the action in the apparatus above referred to is some content that I am confident from my experience that the gas comes over sufficiently pure for ordinary purposes.

Corresiont Method of impregnating Liquids with Sulphydric Acid.



IN Suppose the little flask, F, to contain the liquid to be impregnated, and the flexible pipe, one end of which is inserted into the orifice of the flask, to proceed from a self-regulating reservoir of sulphydric acid: it must be evident that the gas, flowing into the cavity of the flask from the orifice of the pipe, must enter the solution. If not absorbed as rapidly as it may be yielded, the excess must bubble up through the solution; the cork being meanwhile loosened to allow the atmospheric air to escape. The expulsion of the atmospheric air having been completed, and the cork inserted into the neck of the flask, so as to prevent the gas from escaping, it will continue to enter the flask as first as absorbed. But if it be generated in the reservoir more rapidly than the solution can absorb it, the excess must remain in the reservoir, and contribute to depress the acid so low in the bell-glass, as to diminish the quantity of the sulphide on which it can act. Finally, when the solution becomes saturated, the gas generated in the bell must fill it, and thus, by usurping the place of the acid, cause its reaction with the sulplinde of from to be suspended.

'900. Properties.—Sulphydric acid is a permanent ¿ with the odour of rotten eggs, absorbable by water, infla mable and explosive, forming, by combustion with air oxygen gas, water, and a mixture of sulphurous and ¿ phuric acids.

901. At the temperature of 50° F., and under a pression of 17 atmospheres, sulphydric acid becomes a colourk

liquid more fluid even than sulphuric ether.

902. Metals are tarnished by it, especially preparations lead, of which it is a test, and by which it may be detect. It is evolved from privies, blackening the ceruse or carl nate of lead in paint. It may be decomposed by various substances, having an affinity for one or both of its contuents, as for instance, by chlorine, potassium, sodium, sodium, sodium, and ignited carbon; also by successive elements.

tric explosions.

903. Sulphydric acid decomposes all metallic solutions, cept those of cobalt, nickel, iron, zinc, manganese, titaniu and molybdenum, in consequence of the attraction between hydrogen and either oxygen or chlorine, and between metals and sulphur. Metals, which in the metallic st yield hydrogen during their reaction with diluted sulphu or muriatic acid, afford sulphydric acid, when in the st of a sulphide or sulphuret, subjected to those acids. cording to Berzelius, some sulphides act as acids, oth as bases, and unite with each other in a manner analogo to that in which the oxacids and oxybases combine. resulting compounds he calls sulpho-salts. Some sulphi are liable to be reduced by exposure to pure hydrogen a way analogous to that in which oxides are decompo by the same agent. But the number of sulphides wh may be thus decomposed is much smaller. Atmosphe air is said to be rendered deleterious to life by the addit of 110th of this gas.

904. It is alleged that a single cubic inch of the pliberated in a large chamber, will in every part be proceed tive of its characteristic unpleasant odour. A current the gas directed upon the tongue causes an astring acid, and bitter taste. The specific gravity of sulphy acid is 1.1782, that of atmospheric air being 1. I slowly decomposed by nitric oxide, and by sulphurous when moist. Nitroso-nitric acid reacts with it explosive With sulphurous acid when dry it does not react;

eing present, condensation ensues with a deposition ur, and, according to Thompson, the production of ar acid. At the temperature of 50° F., water takes e times its bulk of sulphydric acid, which may be expelled by a boiling heat. The aqueous solution litmus, and becomes turbid after some time by exto the air, with the oxygen of which the hydrogen as combines, while the sulphur precipitates. It has been stated that water impregnated with sulphyid exists in many natural springs, which are much ted by invalids.

The celebrated white sulphur, salt sulphur, and red springs of Virginia, are of this nature. They apurticularly efficacious as remedies for bilious disor-

id in cutaneous diseases.

The red sulphur springs are thought to be pecuseful in some pulmonary complaints, and appear to surprising and unaccountable influence in lowering mency and force of the pulse.

Experimental Illustrations.

Method of extricating sulphydric acid gas by of a self-regulating reservoir exhibited; also, the nation of water with it. Effects of its aqueous soon litmus, and on various metallic solutions. Chawritten with dissolved acetate of lead are blackr exposure to the gas, or its aqueous solution. nation by nitric acid.

Sympathetic Picture.

e original of this figure (see the engraving at the top of the following drawn of a gigantic size, in acetate of lead, and was invisible at a little until a jet of sulphuretted hydrogen was directed upon it. The image wed by the waving of the pipe from which the gas flowed, as if it were of a magician.

the acctate has had time to become dry, the experiment will not succeed storing a due degree of moisture. This object is best accomplished by wet sponge over the back of the sheet on which the figure has been

ienale.—The acetate of lead consists of acetic acid and oxide of lead. a of the oxide unites with the hydrogen of the gas, while the sulphur a sulphuret, to which the blackness of the picture is due.



Of the Polysulphide of Hydrogen.

911. There are various compounds formed by sulphur with metals of which are soluble; as for instance the compound formed by to with lime. This compound has been called a persulphuret of calcium would call it a persulphide. Scheele ascertained that on pouring induted acid a persulphuret, such as that to which I have alluded, a looking liquid was precipitated, which subsequently received the a bisulphuretted hydrogen. Thenard designates this compound as the sulphuret of hydrogen, on account of the great and variable non atoms of sulphur which enter into its composition. Moreover he that it constitutes a compound analogous in its properties to the deut hydrogen; being like that mysterious combination decomposable be substances for which it has no affinity. Even the presence of the phide employed in its production is incompatible with its exister bence the impossibility of forming it by pouring the acid into the s In that case an excess of the persulphide must inevitably be present

912. He alleges that the polysulphuret (polysulphide) is always ordinary temperatures. Its colour is yellow, sometimes approagreenish-brown. It whitens the tongue when applied to it, as is t upon making a similar application of deutoxide of hydrogen. The effect is produced upon the skin. Litmus paper is bleached by a especially when it is diffused in muriatic acid. Sometimes it has a sistency of an essential oil, sometimes of a fat oil, according to the tion of its constituents, which has already been stated to be variated odour is peculiar and disagreeable, especially at the period when been recently formed, the supernatant liquid is decauted from it.

do it affects the eyes painfully. Sooner or later it is resolved into its element spontaneously. Charcoal, platinum, gold, iridium, and many other metals in the pulverulent form, cause the evolution of the hydrogen. Many metallic oxides have the same effect, some so actively as to cause a brist effervescence. These results also ensue from contact with the deutonide or bioxide of manganese, from magnesia, from silica, and above all from pulverized baryta, strontia, lime, potash, and soda. From some of the facts mentioned by Thenard, I infer that this substance may be of great thrice in bleaching.

COMPOUNDS OF HYDROGEN WITH SELENIUM AND TELLURIUM.

Of Selenhydric Acid, commonly called Selenuretted Hydrogen.

913. Selenhydric acid is supposed to consist of one atom of selenium, and one atom of hydrogen. It may be obtained from the selenide of potasun or of iron, by the action of chlorohydric acid. It is a colourless gas, whale without change of colour by water which has been boiled. Water mimpregnated has an hepatic taste, reddens litmus paper, and if applied the skin, stains it a brownish-red. The solution exposed to the air, by exidation of the hydrogen, becomes gradually turbid from the surface wards, acquiring a reddish hue, and depositing selenium in light flocks. I metallic salts, even those of iron and zinc, when they are neutral, are ecipitated by selenhydric acid. The precipitates are generally of a deep ck colour, yet those of zinc, manganese, and cerium, are flesh-coloured. y the oxidizement of the hydrogen in selenhydric acid, selenium is prepitated of a cinnabar-red colour on any moist body. This acid exercises on the respiratory organs a violent action, which might easily become ingerous. It produces at first the odour of sulphydric acid, but soon after prekling sensation in the membranes of the nostrils, which resembles that ented by fluosilicic acid gas, but is more stimulating. Subsequently the se become red, and the sense of smell is paralyzed. A single bubble of tem, received into the nose, caused such a paralysis of the olfactory mes, as to create insensibility even to the fumes of the strongest ammo-The power of detecting odours was not recovered before the expirano of five or six hours.

114. Thenard mentions that Berzelius, in consequence of inhaling sembydric acid gas, was attacked by a cough so severe, that a blister was used secessary. The quantity inhaled was so minute as to give the pression, that, in its effects upon the human system, this gas is presently pernicious.

Of Tellukydric Acid, commonly called Telluretted Hydrogen.

Mhen an alloy of tellurium with zinc or tin is exposed to the action chorohydric acid, telluhydric acid is evolved. It is a colourless gas, it strongly resembles sulphydric acid in smell and in its chemical and chaical properties. It reddens litmus paper, is soluble in water, properties solution, which by exposure to the air becomes brown, the properties of the oxidation of the hydrogen and precipitation of the lit is probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of one atom of hydrogen and one of the probably composed of the p

916. The effect of the monosyllable gen, in chemical language, has be explained. (See note, 628.)

SECTION II.

OF NITROGEN OR AZOTE.

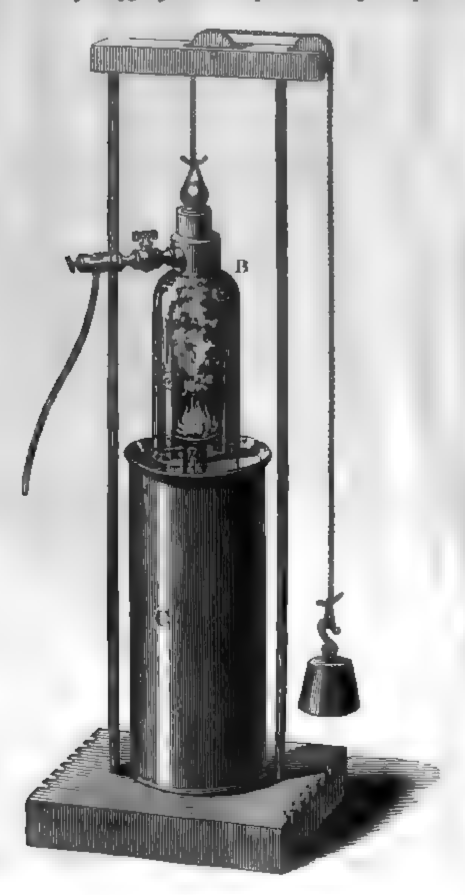
917. In the gaseous state, it forms nearly four-fifths the atmosphere in bulk. Its ponderable base is a princip element in animal substances. In vegetables, it is on occasionally found. It was called azote, from the Gregon, life, and a privative of. It was subsequently naminitrogen, azote being equally applicable to other gas which are destructive of life. I regret that Thenard, i stead of abandoning the use of this bad word, has late endeavoured to give it a further hold on nomenclature, lusing the words azotous and azotic, in lieu of nitrous an nitric.

918. Consistently with the explanation which has be given of the monosyllable gen, nitrogen signifies a capacito produce nitric acid, as oxygen conveys the idea of

capacity to produce acids generally.

- 919. Preparation.—Nitrogen may be procured by 1 aid of any substance which will, in a close vessel, abstration of the atmosphere; a for instance, by the combustion of phosphorus, or by infilings and sulphur moistened. This gas may also be a tained by heating muscular flesh in a retort with nitracid very much diluted. When obtained by means phosphorus, a minute quantity of this substance remain in solution in the nitrogen; when extricated by the action of nitric acid, it contains a small portion of carbonic acid in either case it may be purified by washing it with a alkaline solution, or with lime-water.
- 920. Another method of obtaining nitrogen gas is pass chlorine through liquid ammonia. The chlorine unit with the hydrogen of the ammonia, while the nitrogen liberated. Care must be taken to have the ammonia excess, otherwise a chloride of nitrogen may be form which is capable of producing the most violent exp sions.
 - 921. When the chlorite of lime (bleaching salt) is m

Abstraction of Oxygen from Atmospheric Air by Phosphorus.



(Page 173.)

gled with muriate of ammonia and moistened, nitrogen is evolved. For this purpose Professor Emmet has recommended the boiling of nitrate of ammonia upon zinc.

Apparatus for obtaining the Nitrogen from Atmospheric Air.

22. The apparatus represented in the opposite engraving leaves the nitrogen so stated, as to be drawn easily from the containing vessel, in such quantities and at subtimes as may be desirable. In its principal parts, it does not differ from the someter for oxygen. (617.) It is provided with a pipe, p, concentric with the axis the lower vessel, C, surmounted by a small copper cup. The pipe in question desaids perpendicularly from the level of the brim of the vessel to the bottom; being subtred into a hole in the latter, so that, the bore being accessible from without, the typer cup at the upper end may, when necessary, be touched with the end of a rediction red, introduced through the pipe as in burning phosphorus in oxygen.

23. The inner vessel of the gasometer consists of a bell-glass, B, suspended by a cond pasing over a wooden gallows with suitable pulleys. The bell has a perforated seck remented into a brass cap, furnished with a female screw for receiving a cock. It this cock a flexible lead-pipe is attached by a gallows screw. Upon the copper cap a sufficient quantity of phosphorus being placed, and the lower vessel adequately supplied with water, the bell-glass is suspended within the lower vessel, as is usual with gasometers, and allowed to descend about a third of its depth. Meanwhile, the cock of the tube being open, the air is allowed to escape, so that the liquid within the without the bell-glass may be on a level. The cock being in the next place that, and the temperature of the phosphorus sufficiently raised to make it take fire by tocking the cup with the extremity of an iron rod previously reddened in the lim, a brilliant combustion ensues. As soon as it declines, the iron, meanwhile kept in the fire, should be again introduced, in order to sustain the combustion till all the origen is absorbed.

When the air in the bell-glass is completely deoxidized, which may be known by the funes becoming yellow, the residual nitrogen may be expelled into any recipient at pleasure, through the flexible pipe attached to the cock for that purpose, by

depressing the bell in the water.

925. Properties of Nitrogen Gas.—As a gas, it is distinguished by a comparative want of properties. It is lighter than oxygen gas, or atmospheric air. It supports neither life nor combustion, but is obviously a harmless ingredient in the air.

926. The affinity of nitrogen for caloric, compared with that which it displays for other substances, appears to be peculiarly great. Hence it is not liable, like hydrogen or oxygen, to enter into combination with other matter, so as to part with the caloric to which it owes its existence as a gas; and when under any circumstances it does enter into combination, it seems, more than almost any other substance, to carry caloric into combination with it; being, consequently, an ingredient in a majority of the most powerful fulminating compounds.

927. Nitrogen has been suspected by some chemists to be a compound, but is generally considered as an element. At the temperature of 60° F., 100 cubic inches weigh

30.1650 grains. Its specific gravity, comparatively with air, is 0.9727.

Experimental Illustrations of the Properties of Nitrogen
Gas.

928. A portion of the nitrogen, obtained as above described, being introduced into a bottle, extinguishes a caudle flame when introduced into it; but being mixed with one-fourth of its bulk of oxygen gas, the effect of the mixture in supporting flame is similar to that of atmospheric air.

OF ATMOSPHERIC AIR.

929. Atmospheric air is a mixture, not a chemical conpound, of oxygen and nitrogen gas, with some moistur and carbonic acid, in the following proportions.

	By Measure.						By Weight.	
Nitrogen gas	•	•	•	77.5	•	•	75.55	
Oxygen gas	•	-	•	21.	•	•	23.32	
Aqueous vapour	-	-	•	1.42	•	•	i.03	
Carbonic acid	•	•	•	0.08	•	•	0.10	
				100.00			100.00	

930. The average of a great number of experiment made with my cudiometers, makes the proportion of oxy gen 20.66 in 100 of air.

931. In addition to these constituents, it is alleged that there is a little chlorohydric acid in the atmosphere, is situations in the neighbourhood of the sea; and hence arises, probably, that animals far inland, show a muc greater avidity for common salt, a compound of chlorin and sodium, than those existing in regions bordering of the ocean. This avidity seems to have been implanted if order to supply a source for the chlorohydric acid, whice appears to be requisite to the powers of the gastric fluid.

932. It has been made a question whether the nitroge and oxygen of the air are not in a state of chemical conbination. I am of opinion that no other cause of unic between them exists than that which is known to produt the equable diffusion of heterogeneous gaseous particle among each other, notwithstanding the difference of the specific gravities.

33. In its qualities atmospheric air does not differ from

ixture. Oxygen, mingled with hydrogen in the same portion in which it is mingled with nitrogen in the air, been found to support animal life nearly as well.

4. The mechanical influence of the atmosphere, so far appertains to chemistry, has been sufficiently illusted, (177, &c.) I have also treated of its capacity to moisture, and to promote and produce cold by evaluation (1920, Ac.). Some additional methods of applying tion, (229, &c.) Some additional methods of analyst, will be mentioned under the heads of nitric oxide shosphorus.

Eudiometrical Analysis of the Atmosphere.

i. While on the subject of atmospheric air, the eudiometrical analysis mes necessarily an object of attention. I have already given an sing and description of a large eudiometer, which I have designated rdumescope. By means of that instrument it was demonstrated, that the elements of water are mixed in the gaseous state and ignited, rill always combine in the proportion of two volumes of hydrogen to stame of oxygen. It follows that, if any gaseous mixture containing n, and no other gas capable of combining with hydrogen or oxygen, nted with an excess of hydrogen, all the oxygen will be condensed into , and may be estimated as equal to one-third of the resulting deficit. lows also that, if, to a gaseous mixture containing hydrogen, and no gas with which hydrogen or oxygen can combine, an excess of oxyze added and the mixture ignited, all the hydrogen will be condensed, will in quantity equal two-thirds of the deficit. Thus, if five volumes mespheric air and three of hydrogen be introduced into the volumeand ignited, the eight volumes will be reduced to rather less than five; use a little more than three volumes will have been condensed, of cone-third is oxygen. In five volumes of atmospheric air, there is, bre, somewhat more than one volume of oxygen. By the volumethe excess cannot be accurately measured, but by other instruments I have contrived, and which I shall proceed to describe, great accuis attainable. I am the more particular in describing my apparatus in mpredium, that I may not be under the necessity of occupying the ime allotted to my lectures with such descriptions.

Of the Sliding-rod Eudiometer.

La large constructed some endiometers, and gas measures, in which ensurement of gas is effected by a graduated rod, which slides into or the cavity of a tube, through a collar of leathers soaked in lard, and by a screw so as to be perfectly air-tight. This rod is emto vary the capacity of the tube, and at the same time to be a meafree quantity of air, or of any other gas, consequently drawn in or About one-third of the tube is occupied by the sliding-rod. being recurved, and converging to a perforated apex, is of a form int for withdrawing measured portions of gas from vessels inverted with or mercury.

Libere were two forms of the sliding-rod eudiometer; one designed

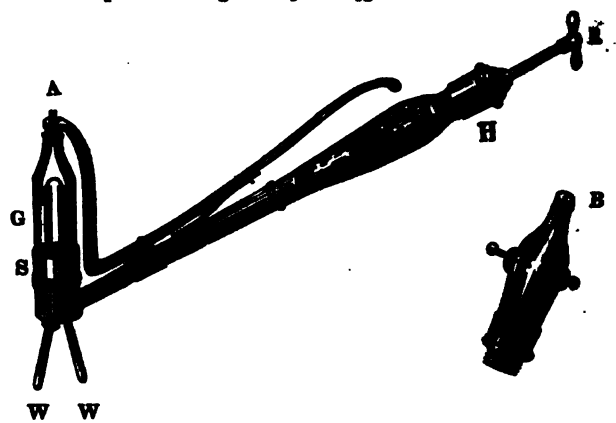
20 Marian Committee

to be used for explosive mixtures, requiring ignition; the other in and dependent upon the absorbing power of a liquid or gas. The former from the eudiometers employed by European chemists, in the contribution igniting the explosive mixtures, as well as in that for measuring galvano ignition (335) being substituted for the electric spark.

938. I shall proceed to describe a sliding-rod eudiometer, for the as sis of explosive mixtures, which I designate as aqueous, because water in

confining liquid employed in it.

Aqueous Sliding-rod Hydro-exygen Eudiometer.



939. This cut represents a hydro-oxygen eudiometer, in which the measures are made by a sliding-rod, and the explosions are effected by the galvanis ign

of a platinum wire.

940. In the instrument represented by the preceding cut, the igniting wire I dered into the summits of the two brass wires, W W, which pass through the be of the socket S, parallel to the axis of the glass recipient, G, within which the seen. One of the wires is soldered to the socket. The other is fastened by a of a collar of leathers, packed by a screw, so that it has no metallic communic with the other wire, except through the filament of platinum, by which they are bly connected above, and which I have already called the igniting wire. The has a capillary orifice at the apex, A, which is closed by means of a lever and a (apparent in the drawing,) excepting when the pressure of the spring is con acted by the thumb of the operator. The sliding-rod, R, is accurately graduat about 160 degrees.

941. Experience has shown the expediency of securing the valve which class aperture in the apex of the instrument from the possibility of leakage during a sions, by means of an iron staple with a screw, represented by the following



This fastens upon two pivots, one of which is inserted on each of the brass socket, S, into which the glass recipient, G, is each ed. The staple hinges upon these pivots, and may be breaght a position in which the screw, A, being immediately over the may be made to tighten it; or the staple may be made to hang a so as not to be in the way when the instrument is to be can In order to use the cudiometer, it must be full of water, free air-bubbles, and previously proved air-tight; the rod being troduced to its hilt, and the capillary orifice open, in consequence of a due degree of pressure on the lever, by which it is not closed. Being thus prepared to ascertain the proportion of each in the air, draw the rod out of the tube till 100 gaduations are ble. A bulk of air, equivalent to the portion of the rod thus

^{*} To prepare the instrument and prove it to be in order, depress the glass ret below the surface of the water in the pneumatic cistern, the capillary orifice

NITROGEN. 177

ill of course enter at the capillary opening; after which the lever must be a close it. Introduce the receiver into a bell glass of hydrogen, and, openwisce, draw out the rod as far as an enlargement upon the end will allow it acted. This arrestation will take place just as the 160th graduation becomes and then, in addition to the 100 measures of air previously taken, 60 of hyll have entered; next close the orifice, and withdraw the instrument from

Apply the projecting wires. W W, severally to the metallic cups, comg with the poles of the calorimotor represented below; then move the as to cause the receptacle holding the acid to rise about the plates. By quent ignition of the wire, the gas will explode. The instrument being rain into the water of the pneumatic cistern, so that the capillary orifice, ed. may be just below the surface, the water will enter and fill up the valed by the condensation of the gases. The residual air being excluded by a portion of the rod remaining without the tube, will be in bulk equivalence, which may consequently be ascertained by inspecting the graduate performed this experiment in thirty seconds.

oxygen is to be examined by hydrogen, or hydrogen by oxygen, we must have a portion of each in vessels over the pneumatic cistern, and successe the requisite quantities of them, and proceed as in the case of atmos-

other and perhaps more accurate mode of operating with this instrument us of one of the volumeters, (see 947, 953,) to make a mixture of the differ, in due proportion, in a bell glass. Thus, let two measures of atmospherided to one of hydrogen; then on taking one hundred and fifty measures ture into the eudiometer, there will be the same quantity of each gas, as ares of hydrogen and 100 of air had been taken, as above described. In scertain the quantity of pure oxygen in the gas from nitre or manganese, are of it might be added to three of hydrogen. Then of 160 measures of re, which might be taken into the eudiometer, 40 would consist of the gas yed, and 120 of hydrogen; and one-third of the deficit, caused by the extould be the quantity of pure oxygen in the 40 measures.

hydrogen were to be assayed, as, for instance, the gas evolved by the reacluted sulpharic acid with zinc, (see page 144,) it would be proper to take to of the hydrogen and oxygen; as the gas which is not to be analyzed are be in excess. Taking then 160 measures into the eudiometer, twothe deficit, caused by the explosion, would be the pure hydrogen in 80 of the gas under analysis. For the last mentioned process it is preferable non the red, in addition to the scale of 160 (942), another of 200 degrees, means fifty measures of oxygen, or 100 measures of hydrogen, may be anathis way the per centage of impurity may be more readily perceived.

tures, in the usual mode for passing the electric spark, should this method against on be deemed desirable, for the sake of varying the experiment, purpose of illustration. This glass, the other being removed, may be fast-the same place. The wires W W, may remain, but should be of such a set to interfere with the passage of the electric spark. The instrument is said usual, excepting the employment of an electrical machine, or electrical the gaseous mixture. For the travelling chemist, the last men-

tand open; draw the rod out of its tube and return it alternately, so that, trake, a portion of water may pass in, and a portion of air may pass out. is operation, the instrument should be occasionally held in such a posture, the air may rise into the glass recipient, without which its expulsion by of the rod is impracticable. Now close the orifice at the apex, A, and a few inches of the rod, in order to see whether any air can enter at the or pass between the collar of leathers and the sliding rod. If the instrument air tight, the bubbles, extricated in consequence of the vacuum promuthdrawing the rod, will disappear when it is restored to its place.

plates of zine and three of copper, severally eight inches by nine, for a description of which, see my Treatise of Galvanism or Voltaic Electricity, writes or cups, in two masses of soft solder which constitute the poles of ment, the wires, W W, of the eudiometer are forcibly pressed by one hand more, while by the other the acid is made to act upon the plates through mentality of the lever. Instantaneously on the ignition taking place, the aid be interrupted by lifting the cudiometer; as otherwise the wire might

tioned mode of ignition may be preferable; because an electrophorus is me

ble than a galvanic apparatus.

946. In damp weather, or in a laboratory where there is a pneumatic c amid the moisture arising from the respiration of a large class, it is often it to accomplish explosions by electricity.

Sliding-rod Gas Measure.

947. The construction of this instrument, represented by the opposite of differs from that of the sliding-rod eudiometers, in having a valve which i and shut by a spring and lever, acting upon a rod passing through a leathers. By means of this valve, any gas drawn into the receiver, is inclu to be free from the possibility of loss, during its transfer from one vessel to This instrument is much larger than the sliding-rod eudiometers for mixtures; being intended to make mixtures of gas, in those cases where be to the other, in a proportion which cannot be conveniently obtained I more or less volumes of the one than of the other, by means of the volumeta 954,) for instance, suppose it were an object to analyze the air according to I son's plan of taking 42 per cent. of hydrogen. The only way of mixing the a volumeter in such a ratio, would be to take the full of the volumeter 21 hydrogen, and 50 times of atmospheric air. By the large sliding-rod gas this object is effected at once, by taking 42 measures of the one, and 100 of the other.

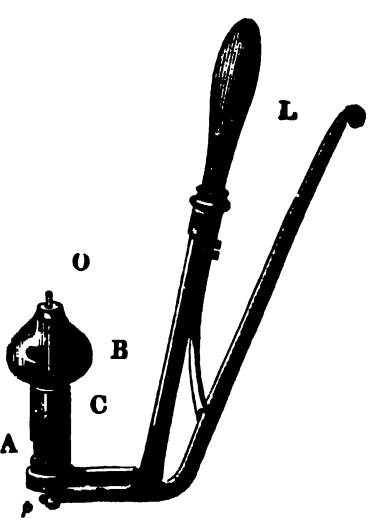
Piston Valve Volumeter.

948. I have contrived some instruments for measuring gas with great acc call them volumeters to avoid circumlocution. They are of two kinds, on by introducing it into any vessel containing the gas with which it is to over water or mercury; the gas is introduced into the other through an ori usual in the case of filling a common bottle over the pneumatic cisternlowing figure will convey a correct idea of one of them, which, having a paralle, I call the piston valve volumeter.

949. The lever, L, is attached by a hinge to a piston, p, which works is chamber, C. The rod of this piston extends beyond the packing through of the bulb, B, to the orifice, O, in its apex, where it supports a valve, this orifice is kept close, so long as the pressure of the spring, acting on

L, is not counteracted by the hand of the operator.

950. Suppose that, while the bulb of this instrument, filled with water or is within a bell glass containing a gas, the lever be pressed towards the har valve consequently is drawn back, so as to open the orifice in the apex of 1 and at the same time the piston descends below the aperture, A, in the



The liquid in the bulb will now cescape, and be replaced by gas, securely included, as soon as the of the spring is allowed to push the beyond the lateral aperture in the ber, and the valve into the orifithe apex of the bulb.

951. The gas, thus included, transferred to any vessel, invendence of the bulb below that of the lever, L, so as the aperture, A, in the chamber orifice of the bulb simultaneously

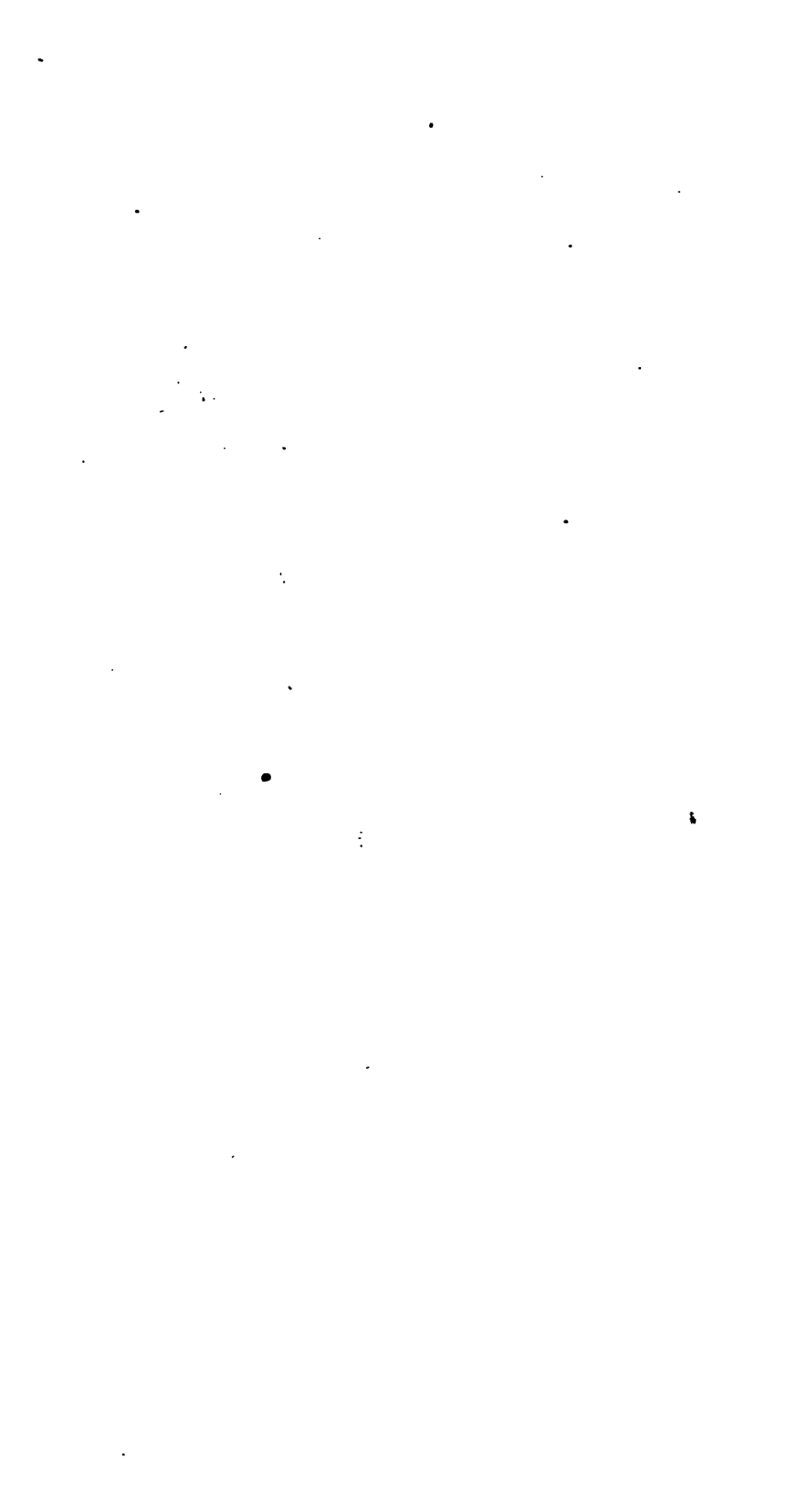
952. The bulk of gas, include volumeter, will always be the sthe quantity will be as the densigns into which it may be introdufilled. Hence, in order to meas accurately, the liquid, whether mercury, over which it may be should be of the same height without. This is especially in the case of mercury, which, weight to water as 13.6 to 1, a density of a gas materially; or

its surface within the containing vessel does not deviate sensibly from the lesurface without.

Sliding Rod Gas Measure.



(Page 178)



Tommove this source of inaccuracy I employ a small gauge, which commu-Livingh a cock in the neck of the bell, with the gas within. In this gauge stiquid will answer, which is not absorbent of the gas. In the case of am-Liquid ammonia may be used; in the case of muriatic acid gas, the liquid The gauge is simply an inverted glass syphon, of one of the legs of which " mile to communicate with that of the receiver, holding the gas, while the gen to the atmosphere. Even mercury may be used in such an instrument afficient accuracy, because the legs of the syphon being near to each other, manute disparity in the heights of the two adjoining columns of the liquid ing the syphon will be discernible.

Simple Valve Volumeter.

954. Besides the lower orifice, O, by which it is filled with gas, the volumeter which this figure represents, has an orifice at its apex, A, closed by a valve attached to a lever. This lever is subjected to a spring, so as to receive the pressure requisite to keep the upper orifice shut, when

no effort is made to open it.

955. When this volumeter is plunged below the surface of the water in a pneumatic cistern, the air being allowed to escape, and the valve then to shut itself under the water, on lifting the vessel it comes up full of the liquid, and will remain so, if the lower orifice be ever so little below the surface of the water in the cistern. Thus situated, it may be filled with hydrogen, proceeding by a tube from a self-regulating reservoir. (797, 798.) If the apex, A, be then placed under any vessel, filled with water and inverted in the usual way, the gas will pass into it as soon as the valve is lifted.

956. Volumes of atmospheric air are taken by the same instrument, simply by lowering it into the water of the cistern, placing the apex under the vessel into which it is to be transferred, and lifting the valve: or preferably by filling it with water, and emptying in some place out of doors, where the atmosphere may be supposed sufficiently pure, and afterwards transferring the air, thus obtained, by opening the valve, while the apex is within the vessel in which its presence is required. In this case, while carrying the volumeter forth and back, the lower orifice must be closed. This object is best effected by a piece of

I, or a pane of glass. It is necessary that the water, the atmosphere, and

hoald be at the same temperature during the process.

IICAL COMPOUNDS OF NITROGEN WITH OXYGEN.

These compounds are five in number, nitrous and wide, and hyponitrous, nitrous, and nitric acid. Their ition is given in the following table:—

Of Protoxide of Nitrogen, or Nitrous Oxide.

This compound does not exist in nature. When obtained it is gaseous; yet the experiments of Mr. Faraday have taught us that under great pressure.

may be converted into a liquid.

959. Preparation.—Nitrous oxide may be obtained the action of dilute nitric acid upon zinc, or by exposi nitric oxide gas to iron filings, sulphites, or other stances attractive of oxygen. It is best procured by su jecting nitrate of ammonia to heat, and receiving the product in an apparatus described in the following article As pure water absorbs this gas, Berzelius receives it eve a saturated solution of common salt.

Apparetus for evolving and collecting Hitrory Online.

960. This apparatus is represented by the opposite engraving A, is a copper value of about eighteen inches in height, and nine inches in domestr, which is represented as being divided longitudinally, in order to show the incide.

The pipe, I proceeds from it obliquely, as nearly from the bottom as possible.

961. Above that part of the cylinder from which the pipe proceeds, there is a disphragm of copper, perfected like a colander. A bell glass is surmounted by a brase cook, C, supporting a tube and hollow ball, from which proceed on opposite asian two pipes, terminating in gallows screws, D D, for the attachment of perfectable brase knobs, soldered to flexible leaden pipes, E E, communicating severally will leathern base. F F, of suitable dimensions.

brass knobs, soldered to flexible leaders pipes, E. E., communicating severally will leathern bags, F. F., of suitable dimensions.

963. The beak of the retort must be long enough to enter the cylinder, so that the gas, in passing from the mouth of the beak, may rise under and be caught by the disphragm. This is made concave on the lower side so as to cause the gas to past through the perforations already mentioned, which are all comprised within a credition in diameter than the bell glam. The gas is, by these means, made to enter the bell glam, and is, previously to its entrance, sufficiently in contact with water, to be purified from the acid vapour which usually accompanies at. On account of the vapour, the employment of a small quantity of water, to want the gas, is absolutely necessary; and for the same reason, it is requisite to have the beak of the retories long as to convey the gas into the water without touching the metal, others with acid vapour would soon corrode the copper of the pipe, B, so us to coable the gas to escape. But while a small quantity of water is necessary, a large quantity a productive of waste, as it absorbs its own bulk of the gas. On this account locative the apparatus here described, in preference to using gasometers or air-holders which require larger quantities of water.

require larger quantities of water.

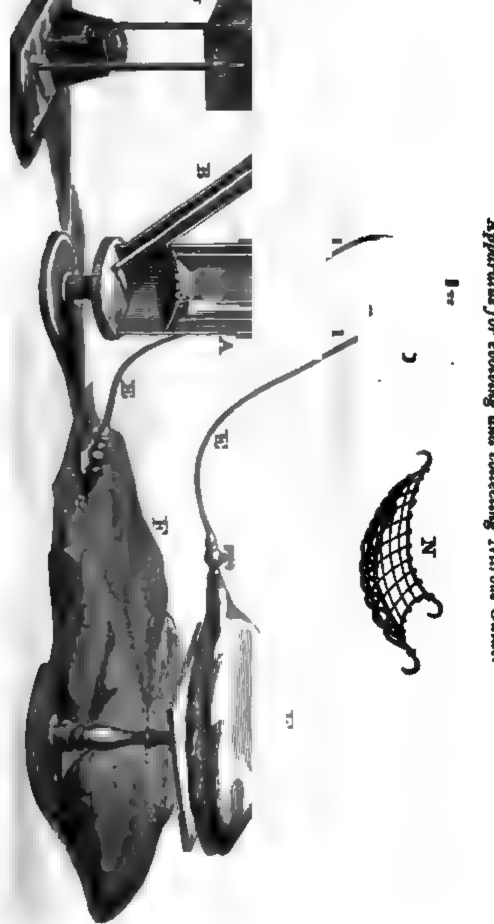
963. The furnace, I, is so contrived, that the coals, being attuated in a draws may be partially or wholly removed in an instant. Hence the operator is some without difficulty, to regulate the duration and degree of the heat. This control of the fire is especially desirable in decomposing the nitrate of aminonia, as the site. otherwise might suddenly become so violent as to burst the relect. The iron settle represented at N, is suspended within the furnace, so as to support the glass term for which purpose it is poculiarly adapted. The first portions of gas which prover, consisting of the air previously in the retort, are allowed to escape through the cock, H. As soon as the nitrous exide is evolved, it may be detected by allowing interpretations of a transfer of a tr

jet from this cock to act upon the flame of a taper.

964. To obtain good nitrous exide gas, it is not necessary that the nitrate of monia should be crystallized; nor does the presence of a minute quantity of main acid interfere with the result. I have employed advantageously, in the products of this gas, the concrete mass formed by saturating strong nitric acid with excess of ammonia.

965. The saturation may be effected in a retort, and the decomposition access plished by exposing the compound thus formed to heat, without further preparation

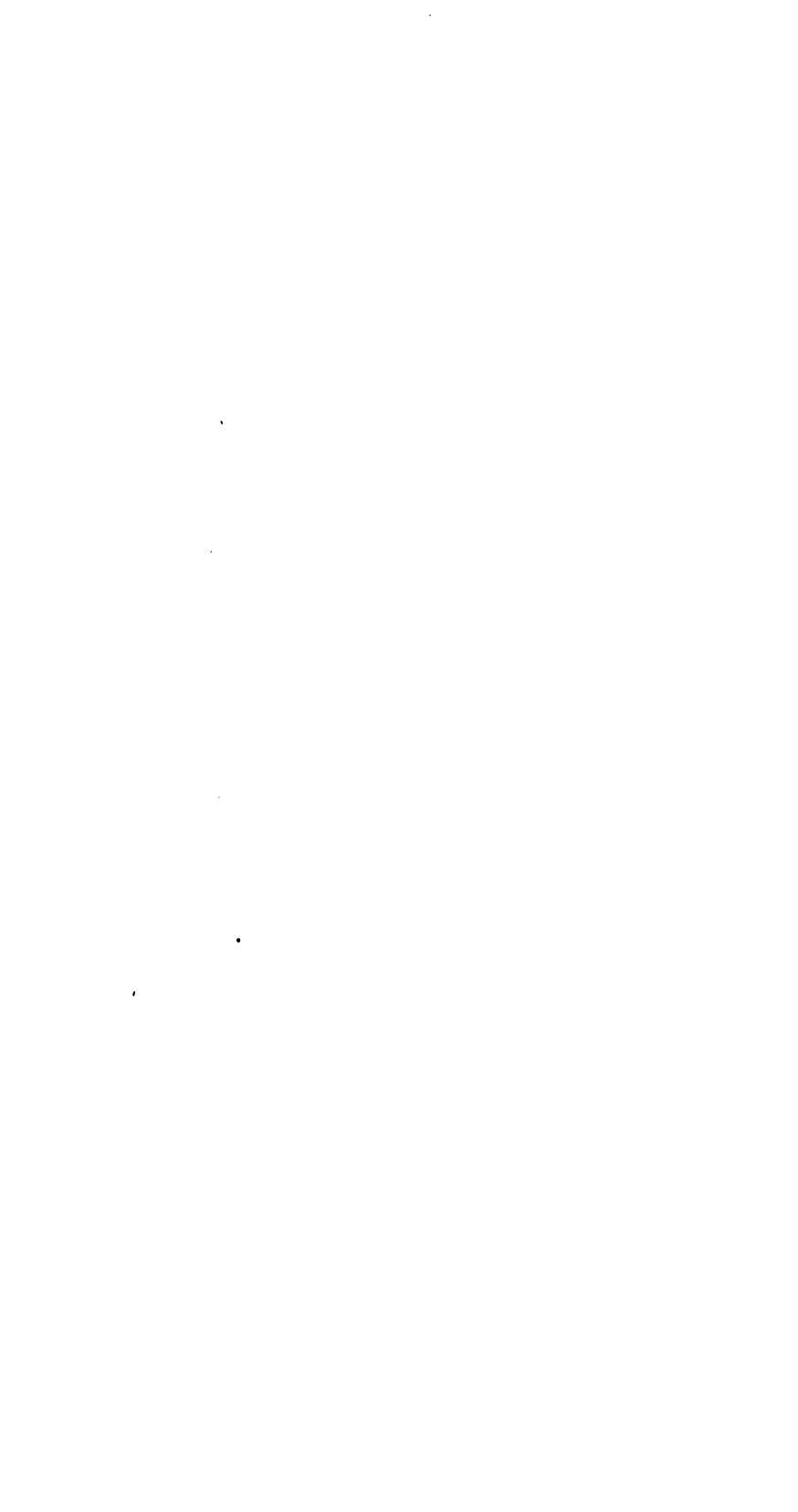
966. Rationale of the Process.—Nitrate of ammonia consists of nitri acid and ammonia; nitric acid, of five atoms of oxygen and one of nite gen; and ammonia, of one atom of nitrogen and three atoms of hydrogen



(Page 180)

Apparatus for evolving and collecting Nitrous Oxide.

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Combustion of Phosphorus in Nitrous Oxide.



(Page 181.)

ne atom of this salt, five atoms of oxygen, three of hydrogen, and two itrogen are, therefore, present. It must be evident that if, in consence of the heat, each atom of hydrogen takes one of oxygen, there will but one atom of oxygen left for each atom of nitrogen. Hence, the le of the salt is resolved into water, and protoxide of nitrogen or nitoxide.

67. Properties of Nitrous Oxide.—It is a permanent . 100 cubic inches weigh 47.25 grains. It supports combustion of a candle flame vividly; though nitric le gas, containing twice as much oxygen, extinguishes ne. Phosphorus is difficult to inflame in it, but burns rapidity when once on fire. The habitudes of sulrare in this respect analogous to those of phospho-. An iron wire burns in it nearly as well as in oxygen . Most of the combustible bodies burn in nitrous oxide. en ignited with hydrogen, an explosive reaction ens, and water and nitrogen result. It has no attribute kidity. When respired it stimulates and then destroys Its effects on the human system, when breathed, are dogous to a transient, peculiar, various, and generally ry vivacious ebriety. It is much more rapidly and exwively soluble in water than oxygen. Homberg's pyroiorus, or that which I have contrived to obtain from ussian blue, takes fire on falling through the gas. Agreeav to the researches of Faraday, to the results of which usion has been made, when nitrate of ammonia was ated at one end of a sealed recurved tube, nitrous oxide w condensed into a liquid at the other end.

Experimental Illustrations.

968. The process and apparatus for producing, collect-& and breathing nitrous oxide gas, exhibited. The efat on a lighted candle and on an iron wire, shown.

Combustion of Phosphorus in Nitrous Oxide.

There is a singular indisposition in the oxides of nitrogen to part their oxygen to phosphorus, until it be intensely ignited either by an execution, or by the access of uncombined oxygen.

This characteristic in the case of nitrous oxide, may be illustrated of an apparatus, like that employed for the combustion of phosin oxygen, and of which the opposite engraving is a representation.

This characteristic in the case of nitrous oxide, may be illustrated to an apparatus, like that employed for the combustion of phosin oxygen, and of which the opposite engraving is a representation.

The upper end of the tube, out-

t cuck, to which a gum chatic

ne receiver having been exhaus previously phoend within the case as the cock communications causes; a illusy entire of the tube, remarking combination, which the mergy.

Of Nitric Oxide, formerly called Nitrous Air-

972. This oxide is an artificial product, and is obtain only in the gaseous state. Its tendency to combin oxygen renders it impossible for it to exist where t mosphere has access.

973. Preparation.—Nitric oxide is evolved during reaction between nitric acid, and copper, silver, and

Mark State

Self-regulating Apparatus for generating Nitric Oxide.



974. The command of a sufficient s nitric oxide is most conveniently atts means of a self-regulating apparatus, the manner which I am about to descr

975. A vessel, perforated at the other respects resembling a decunter; ing a long neck, surmounted by an cap, cock and gallows screw, is place a glass jar of suitable dimensions, a sented in the adjoining figure. By a the gallows screw, a flexible leaden pi attached, as to form a communication bore of the cock. The cavity of th being supplied with copper shreds or t and the jar with diluted nitric acid, by action of the metal with the acid, go piously evolved, producing red fumes l rating nitrous acid with the oxygen of The emission of the gas should be p until the red fumes disappear. The co then be closed, unless it be desirable the gas to be transferred to another ve-

976. It should be understood that the acid passes into and out of the, through the perforation in the stem; while by means of a frag glass, the metallic shreds are prevented from escaping.

977. Properties.—Nitric oxide is colourless, perm ly elastic, and rather heavier than air. By water it slightly absorbed. It is not acid. It extinguishes dle flame, but ignites Homberg's pyrophorus, and su

the combustion of phosphorus, if inflamed before immersion in it, or aided by the access of a minute quantity of oxygen. It is fatal to animals, renders the flame of hydrogen green by mixture, does not explode with it, but explodes with ammonia. It unites rapidly with oxygen gas, the oxygen of the air, or of any other gaseous mixture, producing remarkable red acid fumes. It is absorbed by the green sulphate and the protochloride of iron. The polution acquires the property of absorbing oxygen, and is therefore used in eudiometry. Nitric oxide is decomposed by moistened iron filings; also by ignited charcoal, arse-ic, zinc, and potassium.

Experimental Illustrations of the Properties of Nitric Oxide.

- 978. Copper or silver being subjected to nitric acid, nitric oxide gas is extricated, and collected in bell glasses over water or mercury.
- 979. Absorption of nitric oxide gas by protochloride, and protosulphate, of iron, shown: also the method of ascertaining its purity by the sliding-rod eudiometer; and its application to eudiometry, in various ways, by means of that, and other eudiometrical instruments.
- 980. Self-regulating reservoir of nitric oxide gas for eudiometrical experiments. Absorption of oxygen gas by mixic oxide, and the consequent acidity, made evident by the effect on litmus. Pyrophorus in falling through the gas is ignited.

Of Hyponitrous Acid.

It is acid was isolated in the following manner by Dulong. Having subjected a mixture of four volumes of nitric oxide with one of oxygen, is a tube, to a freezing mixture, he obtained the acid in question in the first of a deep green liquid so volatile as to be converted into a red vapour, the restrained by intense cold. The hyponitrous acid, thus procured, is restrained by water into nitric oxide which escapes; while the composed by water into nitric oxide which escapes; while the composition with another portion of the hyponitrous acid, forms intracid. This unites with the water, and protects the remainder of the hypoirous acid from decomposition. According to Berzelius, it is formed, is combination with bases, when nitrates are kept at a red heat for some line, it is alleged that a hyponitrite of lead is produced, when nitrate of lead is helled with metallic lead.

Hyponitrous acid, when isolated, does not combine directly with the second by contact with them into a nitrate and nitric oxide.

Nevertheless it may be transferred from one base to another. It is

alleged to form a crystalline compound with sulphuric acid, and to combine with nitric acid; but it is questionable whether, in combining with nitr acid, it is not resolved into nitric oxide and nitric acid. in one of the ethereal compounds formed by the reaction of nitric acid wit alcohol. this acid appears to have some practical importance. Of the ethi thus formed it would be premature to treat, until the subject of etherific tion is undertaken.

Of Nitrous Acid.

983. This combination may be procured in the gaseous state, by min ing two volumes of deutoxide of nitrogen, and one of oxygen; or by sul jecting furning nitric acid to heat, and collecting the product in a receive It is also procured by distilling nitrate of lead. Moist nitrous acid is a ge of a deep red colour. When anhydrous, it is a liquid of an orange-yello which boils at 72°. In this form it may be obtained by passing deutoxic of nitrogen and oxygen, both previously dried, through a tube filled wil fragments of porcelain; or by desiccating the nitrate of lead before employ ing it as above mentioned.

984. As the compound, consisting of one atom of nitrogen and for atoms of oxygen, called nitrous acid by the chemists of Great Britain, decomposed when presented to bases. Berzelius does not regard it as a d tinct acid: but gives the name in question, to the trioxide of nitrogen, (756

called hypomitrous acid by the British chemists.

985. The tall receiver and the pear-shaped vessel in its vicinity, being filled wi water, and placed upon the shelf of the hydro-pneumatic cistern, (609, &c.) as repr sented in the engraving, by means of cocks with gallows screws, and a leaden pigproperly attached, render it practicable to make between them a communication

pleasure.

186. These preparations being made, allow the pear-shaped vessel, which I v call a volumeter, to be twice filled with nitric oxide, and as often allowed to yield its contents to the receiver. Then fill the volumeter with oxygen gas. In the n place, open the communication again with the receiver. The oxygen, passing i the nitric oxide, produces dense fumes of nitrous acid. At first, in consequence the rise of temperature which attends the combination, there appears some exp sion; but a speedy absorption of the nitrous acid generated, causes the water to ra and nearly fill the receiver. From some hidden cause, I have never been able attain a complete condensation by this process, however pure might be the gase materials employed.

Application of Nitric Oxide Gas to Eudiometry.

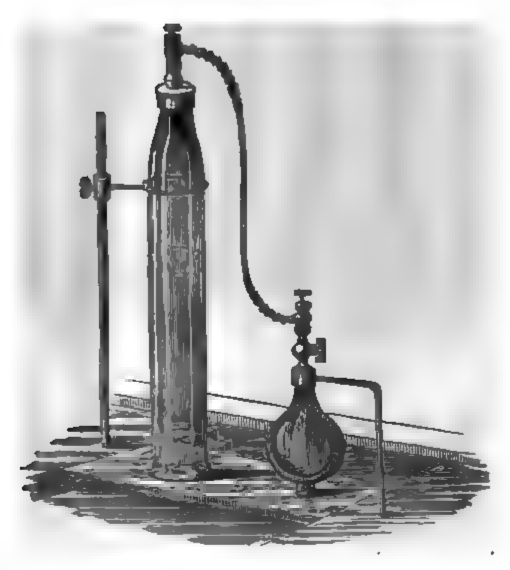
987. The property which this substance has of forming with oxygen, nitrous hyponitrous acid, either of which is absorbed by water, has caused it to be used eudiometrical operations; but owing to the variable proportions in which the above mentioned compounds are liable to be formed, the results obtained have been deemed uncertain, and the directions for using nitrous oxide, given by such em nent chemists as Dalton, Gay-Lussac, and Thomson, are at variance. Gay-Lussi gave an empirical formula, agreeably to which one-fourth of the condensation, po duced by a mixture of equal parts of atmospheric air and nitric oxide, is to be a sumed as the atmospheric oxygen present.

955. As in two volumes of nitric oxide, a volume of nitrogen is combined will one volume of oxygen, occupying the same bulk as if merely mingled,—to conve the nitric oxide into nitrous acid, which consists of the same quantity of nitroge with two volumes of oxygen, one volume of oxygen must be added. Of course, nitrous acid be the product, one-third of the deficit produced, would be the quanti of atmospheric oxygen present. This would be too much to correspond with t

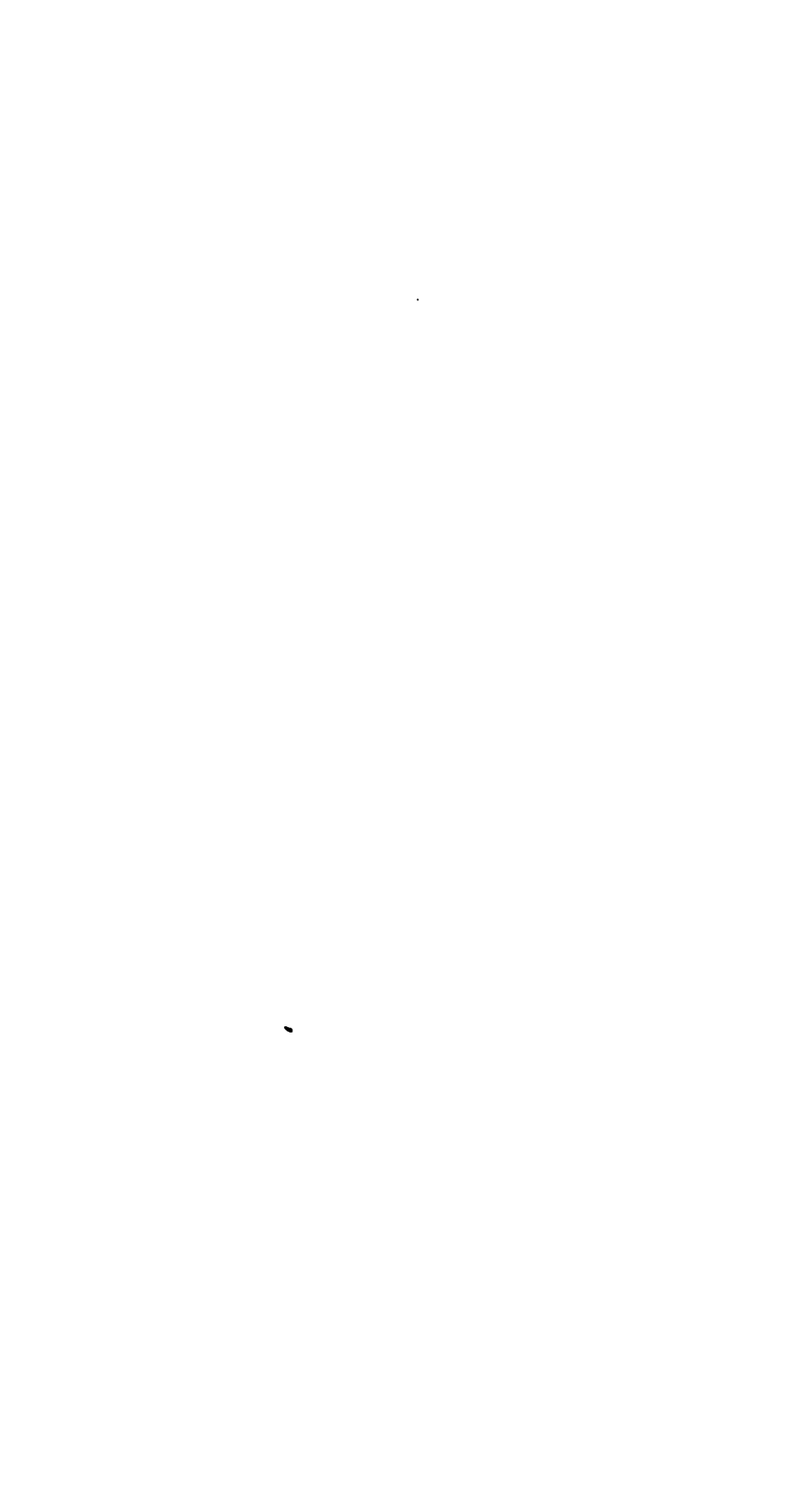
formula of Gay-Lussac.

989. Supposing hyponitrous acid produced, only half as much oxygen would required as is necessary to produce nitrous acid; so that instead of the two volum of nitric oxide taking one volume, they would take only half a volume. The ratio 1/2 in 21/2, is the same as 1 in 5, or one-fifth, which is too little for Gay-Lussac's rul 990. The formula recommended by Dr. Thomson, agreeably to which one-third

Synthesis of Nitrous Acid.

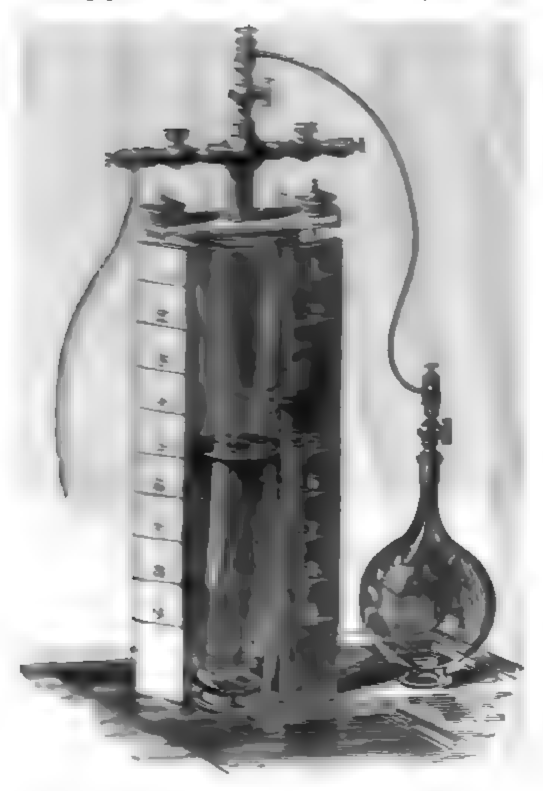


(Page 184.)





Palamenga für die Ameigene of Managéorie Air by Witrie Oxide.



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185 NITROGEN.

eficit is to be ascribed to oxygen, is perfectly consistent with the theory of vos, and much more consonant with the results of my experiments than that renended by the celebrated author of that admirable theory.

I. The late Professor Dana ingeniously reconciled Gay-Lussac's statement with heary of volumes, by suggesting that half a volume of oxygen may take one m of the nitric oxide, and another half volume of oxygen, two volumes.

> do oxygen takes 1 oxide and forms nitrous acid. 2 oxide and forms hyponitrous acid.

The total condensation here would be four volumes, and the deficit due to

n is one volume, or one-fourth.

With the deference due to a chemist so distinguished as the author of the la in question, I long strove unsuccessfully to verify his statements. Agreeably reat number of experiments, annually repeated during many years with differstruments, it has been found that, when three volumes of nitric oxide are mixer water, with five of atmospheric air, nearly the same condensation is effected, en like quantities of air and hydrogen are ignited together. In order to derate the truth of this allegation to my numerous class of pupils, I have emthe apparatus represented by the opposite engraving, and described in the rag article. In this the volumes employed are so large as to make the results wly evident to the most remote observer.

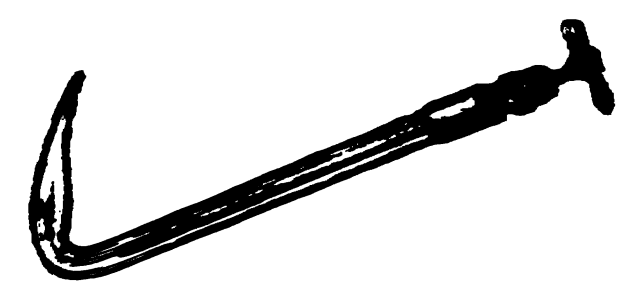
Velumescope for the Analysis of Atmospheric Air by Nitric Oxide.

Secured in a screw rod and plate frame, (248,) there is a glass cylinder thirty in beight, and about five inches in diameter. Into the brass plate which it at top, three cocks are inserted, each provided with a gallows screw. By s of a flexible leaden pipe, let one of the cocks be make to communicate with 'pump. Let the other cock, by like means, be made to communicate with a shaped glass vessel, which acts as a volumeter, or volume measurer. Let the der. by means of scales placed on each side of it, be graduated so as to hold volumes, any three of which shall be equivalent, collectively, to the contents e volumeter The apparatus being thus prepared, and secured over one of the of the pneumatic cistern, (611,) exhaust the cylinder by means of the air pump. to cause the water to rise in it, until by the scale only five volumes of atmosc air are left, and then open a communication with the volumeter. The air ined in this vessel will then pass into the cylinder, consequently the water will to the division upon the scale which designates eight volumes, thus showing recapacity of the volumeter is equivalent to three volumes as premised. Next, mas of the pump, raise the water again to the division upon the scale, marking dames. and fill the volumeter with nitric oxide. If, under these circumstances, mmunication between the pear-shaped vessel and the cylinder be re-established, itric oxide will pass into the cylinder, and, combining with the oxygen of the ined air, will produce nitrous acid in red fumes, which the water will absorb y at first. This absorption is promoted and completed by jets of water, provertically through the mingled gases, by means of the recurved pipe, and lastic bag to which it is attached. It has been shown by the preceding part of scess, that the contents of the volumeter, added to five of air, would make rolumes, were there no absorption; but the actual residue, when the experiis well performed, is always a little less than five volumes, indicating that a sore than one volume of oxygen is contained in the five volumes of air em-. and that this is condensed by combining with twice its bulk of nitric oxide. trous acid, usually thus called, consists of one volume of nitrogen and two m of oxygen. Of course, to convert into this acid, nitric oxide, consisting of dame of nitrogen and one volume of oxygen uncondensed, one volume of amust be added.

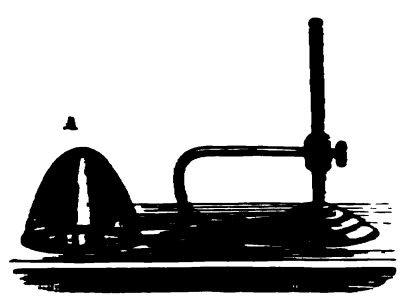
■ Biding-rod Eudiometer for the Analysis of Gascous Mixtures by Absorption.

The form of the sliding-rod eudiometer represented in the next figure does For from that for inflammable mixtures, (940,) as respects the mechanism by the red is secured, or the graduation, which it is convenient to have exactly **both.** The modification which I am about to consider I found very serin the analysis of gaseous mixtures, containing carbonic acid gas, or for

emperaturant de 1937 à l'Alle Clie De 19 le le C. C. Bottomanne d'Antimer des se septent à les califon : le l'amounteme se mile, agressi Le proprie vises : anti-accionis montres.



M. The mexical representation the information, and the transfer of the special states and the second special meaning and the second special states.



A sering filled with water, and immersed in the phenomene circum, the a houng just even with the surface of the water.—In directly out the rod of each conserve, take him the take 10° measures of almostidents are the above described and and have the air in the receiver. While the institute of a set of water, we are any produced from the apex of the instrument, and true the whole of read has gue into the take, communing to have out the rod institute of any appear. In the next place eject the restricting gus from the instrument; we not of graduations of the rod water remain. I the outside if the time, shows the dependenced by the ansorption of the tay gen and nature and in the state of his and

We In a great number of experiments. I have found the deficit to agree nearly with that produced by the explosion of the same quantity of air with a gen, in the aqueous abding-rod hydro-oxygen endiameter, but upon the whole rather greater.

Method of ascertaining the Purity of Nitric Oxide by means of a Solution of Protoculphate of Iron.

We The purity of intric oxide is easily ascertained by means of a solutionation hloride, or green sulphate of iron, and the solution of the salt, and inverted the shelf of the hydro-pneumatic distern, take into the endiometer one hundred nurses of the gas, and transfer them to the bottle, which must be agitated for the minutes. The receiver, being filled with water, and depressed into the of the hydro-pneumatic distern, till the apex. A, is on a level with the surface, up into it the residual gas. In the next place, draw it into the endiometer.

1000. In doing this it is immaterial how much water may follow, because quantity will be inferred from the number of graduations, which must enter the of the tube, in order to effect the expulsion. Of course the impurity will be number thus found

A starated solution of nitrie oxide in the abovementioned ferruginous soluties power of absorbing oxygen, and was recommended by Sir H. Davy as a secretaining the quantity of that gas in the air. The mode of using the the same as that just described, taking oxygen into the endiometer intric oxide, and filling the bottle with the ferruginous solution of nitric tend of the solution of the pure sulphate or protochloride of iron. I have such less satisfactory than those already described.

Theory of Volumes.

Lit is presumed that a reader, who has carefully this work thus far, may have his attention advantly directed to the theory of volumes; otherwise guage, now usually employed in treating of combinesulting from the union of gaseous substances, not always be intelligible to him.

It has been advanced by Gay-Lussac, that sub-, when aëriform, unite in volumes which are equal, when unequal, the larger volume is double, triple,

iruple the other.

This hypothesis has been verified by experiment espect to all substances which are capable, while s, of being combined or decomposed. It is extended exerce to other substances, under the idea that all ceptible of the aëriform state. A volume is said to equivalent of another volume, when capable of y with it a definite compound, or when just adeo displace it from combination.

It must be evident, a priori, that if each atom, of er kind, were to occupy in the aëriform state an space, atoms might be as well represented by equal as by their equivalent numbers; the former after measure, what the latter give by weight. Now mee justifies the belief that, in general, atoms do an equality of volume when rendered aëriform, and then the bulks assumed are unequal, the inequality e removed by multiplying or dividing, by a whole those volumes which are smaller or larger than the This is all that the hypothesis of Gay-Lossac

Berzelius infers that water, and the protoxides of and nitrogen, each consist of one atom of oxygen, atoms of the other ingredient. Admitting this to meet inference, equivalent weights of the four elegaseous substances abovementioned, actually oc-

cupy equal spaces; so that their atoms are as well a sented by equal volumes, as by the numbers indic their ratio to each other in weight. But if we say that, in the compounds abovementioned, there is only atom of each ingredient, the equivalent volumes of rine, hydrogen, and nitrogen, although still equal to other in bulk, will each be twice as large as the equivolume of oxygen. The British chemists, in general ferring the last mentioned view of the atomic constitution of the compounds abovementioned, represent the atomic chlorine, hydrogen, and nitrogen, each by one volume atom of oxygen by half a volume.

1007. When gaseous substances enter into combine preserving the aëriform state, in some cases there is duction of volume, in others none. When a reduction of volume, in others none. When a reduction of the bulk, or resulting volume of the compais to the aggregate bulk of the constituent volumes, as 1 to 2, 1 to 3, 1 to 4, or 2 to 3, 2 to 5, &c.

1008. This will be rendered evident by the foliotable, in which the number of atoms and the number volumes which enter into some important compared are represented by corresponding squares. Each a stands for a volume, and half a square for half a volume first column contains the name of each gas a pour; the second, the equivalent volume of the gas if ple, or an association of the volumes representing constituents if compound; the third, the resulting volume of the compound formed; and the last, the pressur pressed in atmospheres, necessary to produce liquesary.

1009. Among the instances cited in the table, it v seen that there is none in which the bulk of the c tuent volumes is to that of the resulting volume in a greater than that of 3 to 1. The only permanent which the elements are alleged to exist in a stagreater condensation, is olefiant gas, consisting o volumes of the vapour of carbon, and two volumes of drogen, condensed into one volume. There are vapours, consisting of the same elements in the atomic proportion, in which 8 or 9, or, according to Thomson, even 25 constituent volumes are container resulting volume.

MITROGEN.

Tails of the Equivalent Weights and Volumes of some Gases and Vapours.

		1	Pressure of
ited Vepours.	Composent Volumes.	Resulting Volumes of Compounds.	Liquefaction Atmospheres.
	08		
ine	C1 36	<u>.</u>	4 at 60°
ziù of chio-	Cl 36 O8	44	
gen	H)1	
	H 08	9	
obydnic acid	C H 36 1	37	40 at 50°
 .	N 14		
agheric air .	N 14 N 14 O 8	36	
m ozide	N 14 08	22	50 at 45°
ozi de	N 08	30	
≅ eid	$ \begin{array}{ c c } \hline 08 \\ \hline 08 \\ \hline 08 \\ \hline 14 \\ \hline 08 \\ \hline 08 \\ \hline \end{array} $	46	
.	H 1	17	6½ at 50°
t	N H 1		

Of Nitric Acid.

will not combine with oxygen; yet, when mixed with and exposed to a succession of electric sparks, nitric acone of the most important agents in chemistry, is generated. Berzelius alleges that traces of nitric acid, in combination with ammonia, may almost always be discovered the rain water accompanying thunder storms. This chemical combination is probably produced by lightning. The same author states that when a jet, consisting of one while flowing into a vessel containing oxygen, nitric acid, in combination is probably produced by lightning. The same author states that when a jet, consisting of one while flowing into a vessel containing oxygen, nitric acid, in combination is probably some unknown means by which chemical union is induced between nitrogen oxygen; whence the great quantity of nitrate of potential produced in various situations.

1011. It has been supposed that this acid is form during the eudiometrical analysis of atmospheric air hydrogen; and that the deficit being thus increased to an undue estimate of the oxygen. I consider this pression erroneous; as upon one occasion, by exploding successive portions of hydrogen with atmospheric air, collected nearly half an ounce of water, and found it developed.

of acidity.

1012. Preparation.—The production of nitric acid be electricity is too laborious to be resorted to for the purpos of the chemist.

sists of nitric acid and potash, is subjected to heat with a equal weight of sulphuric acid, in a glass, porcelain, or incretort, communicating with a glass receiver. The nitracid is displaced by the superior affinity of the sulphur acid for the potash, and, being vaporized by the heat passes into the receiver, where it condenses into a liqui. Thus obtained, it is more or less contaminated with nitrous acid or nitric oxide, also with chlorohydric, and sulphur acid. By distilling from it about a third of the who quantity, the nitrous and chlorohydric acids pass over in the receiver with the portion of nitric acid distilled, leaving the residue in the retort free from them. Sulphuric at may be removed by distilling the nitric acid from of eighth of its weight of pure nitre, or by the addition of the sistemans.

which precipitates in the form of an insoluble comwith the sulphuric acid. Chlorohydric acid may in unner be removed by a solution of silver; as this orms with the chlorine an insoluble compound which tates.

. Properties.—Nitric acid emits pungent suffocating and has a peculiar odour. When pure it is colourst when exposed to the light, it is slowly decomnto oxygen gas, and nitrous acid or nitric oxide is absorbed, giving an orange colour to the nitric This decomposition takes place much more rapidly un. Nitric acid cannot be obtained free from wa-Vith almost all the metals it reacts powerfully, also ganic substances, causing them to be oxidized. It and destroys the skin. It may be considered as ng of the ingredients of atmospheric air in the liquid ut containing ten times as much of the active prinxygen. It is the most energetic principle in gun-In its highest state of concentration, at a specirity of 1.55, one atom of the acid contains one atom This concentrated acid boils at 175° and freezes P. When it contains one atom of acid to four of it has a specific gravity of 1.42, and boils at 248°. cid, whether weaker or stronger than this, has the point at a lower temperature. If weaker it is bened, if stronger it is weakened, by boiling; and f all degrees of strength become, by the continued tion of a sufficient degree of heat, of the specific of 1.42. The officinal specific gravity is 1.5, in case it contains two atoms of water to one of acid. i. Nitric acid is employed in giving a yellow colour, various other puposes in manufactures. It is used icine for fumigations, in cases in which chlorine is ble.

Experimental Illustrations.

by means of a glass retort and receiver, heated by or small sand bath. Its action on various sub-

Of the Orange-coloured Fuming Nitric Acid, called Nitrie nitric Acid in the Swedish Pharmacopaia.

1017. In whatever proportion sulphuric acid may be apployed in the process just described for procuring nits acid, the liquid obtained is of an orange colour. lour becomes deeper, when the quantity of sulphuric aci employed is insufficient to produce a bisulphate with the potash. I am under the impression that, in some degree the same result follows when the acid exceeds the propor tion requisite to produce the bisulphate. In either case the water, which in the absence of some other base is indispen sable to the existence of nitric acid, is not furnished in sufficient quantity. Hence the acid is partially resolve into oxygen and nitrous acid, which latter, together will the nitric acid, passes into t receiver, constituting t orange-coloured fuming liqu, mentioned by Berzell under the name of nitroso-nit acid. This acid, by expe sure to heat, disengages nitrous acid gas, and becomes co lourless nitric acid. Nitroso itric acid ignites countie oils, carbon, and phosphorus; the latter explosively. It i much more energetic in its reaction with such substance than pure nitric acid, which, probably, when nitric oxide i not present, requires for its existence a larger proportion of water. I deem it probable that it is with nitric oxide not nitrous or hyponitrous acid, that nitric acid is com bined in nitroso-nitric acid. Berzelius conceives tha either view of its composition may be correct.

Experimental Illustrations.

1018. Reaction of nitroso-nitric acid with carbon and oil of turpentine, exhibited, also with caoutchouc tar.

Of the Agency of Nitric Oxide in generating Sulphuric Acid.

vapour are mingled, a crystalline compound is formed, which, if the opt ration be performed within a glass vessel, will appear upon the interior set face in a 'crystalline deposition, resembling hoar frost. When water I added to this compound, it is resolved into sulphuric acid and nitric oxide. The former combines with the water, while the latter escapes in the gasest form, producing with oxygen, if present, the red fumes of nitrous, or hyperitrous acid. It may be inferred that hyponitrous acid, produced as about mentioned, yields one atom of oxygen to the sulphurous acid, converting into sulphuric acid. The acid, thus produced, unites with the nitric oxide.

water: but on being subjected to a larger portion of water, for which has a greater affinity, the nitric oxide is allowed to escape. These habilies of the agents in question excite greater interest, on account of their sency in the generation of sulphuric acid, one of the most valuable of the seruments which have been placed within the reach of the chemist, artist and manufacturer.

Experimental Illustration of the Reactions which occur in the Manufacture of Sulphuric Acid.

1020. Into a glass globe with three tubulures, insert through one of them, t bak of a pint retort, containing about a pound of mercury, and as wh sulphuric acid as will cover it to the depth of half an inch, applying in retort a chauffer of coals. Into the other tubulure, fasten the termicon of a pipe proceeding from a self-regulating reservoir of nitric oxide • The third tubulure should be closed by a glass stopple. The mercury some atom of oxygen from the sulphuric acid, converting it into suland acid which enters the globe. As soon as this appears to have taken ce, a portion of the nitric oxide gas is allowed to enter from the opposite e. Meeting with atmospheric air within the vessel, the nitric oxide will duce red fumes, which, encountering the sulphurous acid, will condense o a crystalline deposition. Occasionally, the stopple must be lifted to bw the access of fresh air; and the supply of this and the gases must so regulated, that the red fumes shall be repeatedly produced and con-When a deposition of crystalline matter, sufficiently striking, has en produced, if water be poured into the globe, the deposition will be wdily decomposed with an evolution of nitric oxide. This gas, meeting ith the oxygen of the air, produces red fumes, which, by the readmission sulphurous acid, are again condensed with it into crystals. These crys-- as before, by the addition of water, are decomposed into nitric oxide s and sulphuric acid. The water in the globe, being decanted and tested, vs decided indications of the presence of sulphuric acid.

1021. Latterly, the process above described, has been resorted to in the large way in the manufacture of sulphuric acid. In some cases the translate has been evolved by the reaction of nitric acid with organic the large of a nature to produce oxalic acid, but in other manufactories relate oxide is obtained from nitric acid by subjecting it to sulphuric aid, who a causes it to be resolved into nitrous or hyponitrous acid fumes of largen gas.

1922. When pitric oxide is obtained by the reaction of nitric acid with war molasses, oxalic acid is produced, and tends to defray, partially, warmse of the process.

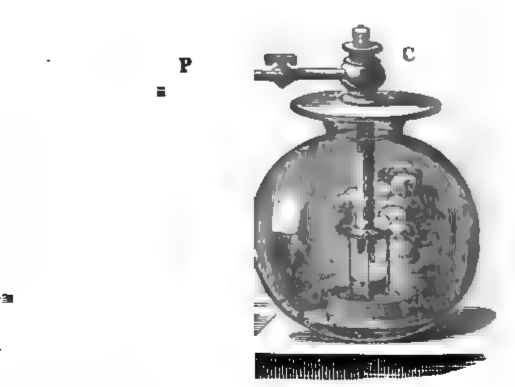
Of the Process usually employed in the Manufac are of Sulphuric Acid.

The combustion of one portion of sulphur, and the simultaneous deflagrament in the mire of another portion, (the fumes created in both ways being received ingrechamber lined with lead, and covered at bottom with water) are the means only employed for the manufacture of sulphuric acid. Each atom of nitre control at atom of potash and an atom of nitric acid. Three out of the five atoms regress meach atom of the acid, unite with an atom of sulphur, converting it into phone acid, which combines with the potash. The two remaining atoms of oxystemetric with the introgen of the acid, are evolved as nitric oxide, which, with rephenic oxygen, moisture, and the sulphurous acid produced by the burning that, generates the crystalline compound above described. Of late years, the two senses of an adequate quantity of moisture has been insured by the introduction

of steam at proper intervals. The crystalline compound, subsiding into the wais decomposed into sulphuric acid, which remains in solution, and nitric oxide, oxide, meeting with further portions of oxygen and sulphurous acid, again or butes to the formation of the crystalline compound, to be again decomposed, process is continued until the water in the chamber becomes sufficiently improgravith sulphuric acid, when it is transferred to leaden boilers. In these it is out trated by boiling, but it is removed before it attains sufficient strength to attack lead, to a platinum alembic, or to glass retorts, and boiled down to the specific vity of 1.85. After it has reached that density, no farther concentration can be facted by heat. This, accordingly, is the standard specific gravity of the sulphacid of commerce.

Production of Sulphuric Acid further illustrated.

1024. The apparatus here described, serves to show, in miniature, the process generating sulphuric acid.



1025. Provide a globular glass vessel with a wide mouth fitted to a suitable cand capable of holding at least eight gallons, represented by the preceding fit Through a hole in the centre of this plate, a gun barrel, open at both extremitionade to descend. From the lower extremity, a ring of about two inches in diter is suspended by wires, hooked to a perforated circular piece of sheet metal, we encircles and is soldered to the barrel. In the ring thus suspended, a conical turn of iron, having an hemispherical cavity, is scated, so as to be a little above water. Between the outside of the gun barrel, and the inside of the brase cast, which supports it, there is a passage from the pipe, P, into the cavity of the galass. A tube leads from a suction pump into this vessel, which is placed of shelf of the preumatic cistern, covered with water as usual.

1626. The apparatus being thus arranged, the inetallic plate, with the gun bring, and frustum appended to it, must be removed from the globe, the iron frushfield out of the ring, and some nitrate of potash (mitre) being introduced intervity in the frustum, it must be made moderately red-bot. It is then to be rest to its seat in the ring, below the gun barrel, and the plate and gun barrel mureturned to their previous position over the mouth of the globe, so that the amay be situated as represented in the engraving. Lumps of brimstone, about size of peas, are to be dropped through the gun barrel into the melted nitre each lump reaches the nitre a combustion ensues, equally remarkable for beaut brilliancy. The globe then becomes filled with sulphurous acid gus, accomp by nitrie oxide gas, and a crystalline deposition ensues. Meanwhile, to keep supply of oxygen within the globe, and to prevent the escape of furnes integration, the suction pump is put into operation, in order to draw the furness the globe, and cause them to be replaced by air, which enters through the guilties are the globe, and cause them to be replaced by air, which enters through the guilties are the globe, and cause them to be replaced by air, which enters through the guilties are the globe, and cause them to be replaced by air, which enters through the guilties are the globe, and cause them to be replaced by air, which enters through the guilties are the globe.

The water rises from the cistern into the bell, until the resistance which it this to father elevation, is greater than that which the water, in the tumbler on the stand, opposes to the entrance of air from the pipe; and, consequently, the air is then the globe through the water in the tumbler, by which the fumes arising the combustion are arrested, especially if liquid ammonia shall have been pre-

1627. To protect the globe from the heat of the red-hot iron frustum, a cylinder of

that kad is placed below it, as represented in the figure.

It is fumes generated during this process, condense upon the inner surface the globe, into a white crystalline compound, identical with that procured in the second above described. By the affusion of water, this crystalline matter undergoes accomposition like that already described, (1020,) giving out nitric oxide, and yield a sulphuric acid to the water.

COMPOUNDS OF NITROGEN WITH CHLORINE AND IODINE.

1029. Neither chlorine nor iodine combines directly with nitrogen; but but unite with the nitrogen of ammonia, under circumstances which I shall mention presently.

Of Chloride of Nitrogen.

1030. This compound may be obtained by placing a bell glass, filled with chlorine, over a solution of one part of nitrate of ammonia in twelve of water, at the temperature of 70°. The chloride appears in drops, which remble olive oil, and which, being heavier than water, subside to the bottom of the basin containing the solution. It is remarkable that this substance does not explode with many combustibles, which would appear more likely to decompose it than those with which it does explode. Thus it explodes with turpentine or caoutchouc, but not with camphor.

1031. The force with which a minute portion of chloride of nitrogen explodes, on contact with oil of turpentine, would hardly be credited by those who have not witnessed this phenomenon. An open saucer of Canton china was fractured by a globule not larger than a grain of mustard seed. The glass tube employed to project the globule into the saucer, was violent-

ly dispersed in fragments.

Of Iodide of Nitrogen.

1032. When iodine is kept in liquid ammonia, it is converted into a brownish-black substance, which is iodide of nitrogen, and which may be collected and dried on bibulous paper at a gentle heat. The iodide of nitrogen thus formed, evaporates spontaneously. It explodes by a slight presure, or when heated or much dried, being resolved into nitrogen gas and iodine.

ON SOME POINTS OF CHEMICAL THEORY.

logs of the phenomena of combustion, and of the properties of some acids, to render it expedient to present to him some general views of combustion, existly, and alkalinity, and additional instruction on classification and nemacleture. I am the more inclined to this course, as, among the compounds of nitrogen, there are three acids and an alkali.

Of Theories of Combustion.

1034. Stahl supposed the existence, in all combustibles, of a caprinciple of inflammability, which he called phlogiston, from pays burn. He inferred that all substances, in burning, gave out phloging combustion, obviously in consequence of the absorption of oxygen the atmosphere. By the advocates of the phlogistic theory, nitrogen confounded with carbonic acid, and carbon with hydrogen, because carbon and hydrogen were conceived to consist of phlogiston nearly and oxygen, in combining with them, was supposed to become plicated air, the name then given to nitrogen gas. It is now well know with carbon, oxygen forms carbonic acid, with hydrogen water; as nitrogen gas contains neither carbon nor hydrogen.

1035. Sulphuric, and phosphoric, acid, and metallic oxides, west rally supposed to be ingredients in the sulphur, phosphorus, and producing them. Thus of two bodies, that which was actually the

was assumed to contain the other.

1086. The celebrated Lavoisier, to whom we are chiefly indebted a exposure of these fallacies of the theory of phlogiston, having ascer that oxygen is an indispensable agent in all ordinary cases of combo was erroneously led to infer that it was in all cases necessary to the cess. But it is now well known that there are many instances of cutton, in which oxygen has no agency.

1037. I would define combustion to be a state of intense corpused

action, accompanied by an evolution of heat and light.

1038. That increase or diminution of temperature consequent to a cal combination, which constitutes combustion when productive of he light, has been ascribed to a mysterious law, by which bodies und change in their capacity to hold caloric. It has been supposed that a pacity of the compound is in some instances greater, in others less the mean capacity of the constituents; and that in the former case un followed by an absorption of caloric, and of course by cold; in the by the expulsion of caloric, and, consequently, the production of Yet, when the capacities of compounds are compared with those of the gredients, the result does not justify the idea that the heat given out latter in combining, is produced by a diminution of capacity. At be hypothesis only substitutes one enigma for another; since it does a count for the alleged change of capacity.

1039. The diversity of power to hold caloric in a latent state, tech designated by the word capacity, is now generally ascribed to the inting influence of electricity. It has been shown* that, if neighbouri dies be electrified by means either of glass or resin, previously subjective friction, they will repel each other; but that if one be thus excited by and another by resin, attraction between them will ensue. Hence citements are considered of an opposite nature. It will be recollected according to the Franklinian theory, the vitreous excitement results redundancy; the resinous, from a deficiency of the electrical fluid former being designated as positive, the latter as negative electricity, ably to the doctrine of Dufay, the different electric excitements are dered as the effects of two different fluids, attractive of each other, b

^{*} See my Treatise on Statical Electricity.

mellent. The one has accordingly been called resinous, the other vitreous dectricity. Yet, even by electricians, who suppose the existence of two faids, the terms positive and negative are employed.

1040. It has been suggested that Voltaic phenomena, combustion, acidity, alkalinity, and chemical affinity, may owe their existence to the principle by which the different electric excitements are sustained in electrified bodies, modified in some inexplicable manner, so as to act between atoms instead masses. This suggestion derives strength from the following facts, which

bee been fully illustrated in my lectures on electricity and galvanism.

1041. The pole of a Voltaic series, terminated by the more oxidizable mani, has been shown to display a feeble electrical excitement, of the same had as that which is producible by friction in glass; while the other pole diplays the opposite excitement, in like manner producible in resin. From reserated experimental observation it is now generally inferred, that, of any two elementary atoms, chemically combined, and simultaneously exposed, to the voltaic current, one will go to the positive, the other to the negative pole. Atoms are supposed to have electrical states the opposite of those of the poles at which they may be liberated, and are said to be electro-negative when liberated at the positive pole, or anode; electro-positive when liberated at the negative pole, or cathode. See my Treatise on Galvanism, page 7.

1042. Substances which have opposite relations to the Voltaic poles, have mathrity for each other, which is usually stronger in proportion as the twenty of their electric habitudes is the more marked. Thus, for interest, oxygen, which is pre-eminently electro-negative, and potassium which is pre-eminently electro-positive, have, under ordinary circumstances, a pre-

dominant affinity for each other.

1043. On all sides it must be admitted that between chemical reaction, raivanism, and electro-magnetism, there is an intimate association which must be explained before the phenomena of chemical reaction can be well understood.

1044. It has been mentioned that, of known bodies, oxygen appears to be most electro-negative. It is questionable whether the grade next to ayen, in the electro-negative scale, is to be assigned to chlorine or fluorine. The follow bromine, iodine, sulphur, selenium, and tellurium.

1045. Among the metals we have a series of substances, varying from the in which the electro-positive power is pre-eminently great, as in potaswan sodium, lithium, barium, calcium, magnesium, &c., to such metals as belong rather to the electro-negative class. Hence, setting out from the streme abovementioned, we may proceed through a long range of metals and less electro-positive, till we arrive at such as produce electro-Entire combinations with oxygen or chlorine, or both. More or less within this predicament, I think we find tin, mercury, gold, platinum, pallam, antimony, arsenic, molybdenum, and lastly tellurium. Thus at an wrom-nate point between the extremes at which oxygen and the alkalifiwhich are placed, there are substances whose relation to the Voltaic pois sequivoral or wavering; and it should be understood that this relation Chlorine is electro-positive with oxygen and persalvave comparative. haps burnne, and electro-negative with every other body. lodine is electropositive with oxygen, chlorine, bromine, and probably fluorine, while with other substances it is electro-negative.

^{*} See my "Treatise on Galvanism, or Voltaic Electricity, for Effects of Galvanic voltair Circuits." page 19. And for Theory of the same, page 35.

1046. Substances of the two opposite classes, in combining with other, constitute compounds which are either electro-positive or elemegative, accordingly as the different energies of their ingredients prederate. Thus in alkalies, consisting of oxygen united with the alkalid metals, the electro-positive influence predominates; while the reverse is of acids, consisting of the same electro-negative principle, oxygen, in bination with sulphur, nitrogen, phosphorus, carbon, boron, silicon, nium, or other substances, which, in their electrical habitudes, lie bet oxygen and those metals.

1047. In some cases we see an electro-negative or electro-positive p attached to compounds, which is not equally displayed by either of constituent elements separately. Cyanogen, consisting of carbon an trogen, is a striking instance of an electro-negative compound thus co tuted; and in ammonia, and the vegetable alkalies lately discovered have instances of electro-positive compounds, produced from principles.

paratively electro-negative.

1048. For any further view of the connexion between chemical galvanic reaction, I refer to my Treatise on Galvanism, or Voltaic Ekcity, especially to pages 7, 17, 85.

Of the Influence on Classification and Nomenclature of the Habitud Chemical Agents with the Voltaic Series.

1049. It would follow from the statements made under the last I that there should be a resemblance between the properties of substational which have a proximity to each other in the electric series. (1042.) cordingly we find, that those which occupy the higher part of the denegative scale, have, by distinguished writers, especially in Great Bribeen classed as supporters; while those which are electro-positive, or followed electro-negative, have been by the same authors classed as combustional Also, certain electro-negative compounds, formed of the pre-emine electro-negative principles, have been associated as acids; while other pounds, of oxygen at least, which have the opposite polarity, have associated as bases, under some of the subordinate divisions of alkaline earths, earths proper, or simply oxides.

1050. The idea of a class of supporters of combustion, and of combust has no better foundation than that certain substances are the most free agents in combustion. Thus hydrogen will produce fire with oxygen chlorine only; sulphur with oxygen, chlorine, and the metals; and bon with oxygen; but as either oxygen or chlorine will burn with a gr variety of substances, they have been called supporters of combustion the substances with which they combine during combustion, combust Iodine and latterly bromine have been classed among the supporters cause they combine with almost all the bodies with which the other ments classed under that name unite, and in some cases with an evol of heat and light. Yet they are not gaseous like oxygen and chlorine are as analogous to sulphur as to oxygen. There appears to me to error in taking either of these substances into the class of supporters, sulphur is excluded, which, next to oxygen and chlorine, has the pro of burning with the greatest number of substances. In other respect phur seems, in its properties, to be intermediate between iodine and The habitudes of selenium appear to range between those c lurium and sulphur.

1051. Hydrogen, phosphorus, carbon, boron, and silicon are no mo

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se called comb so, than oxygen, chlorine, bromine, and iodine, a called supporters. It should be observed, also, that these appellatived in hydrogen, is productive of a flame, similar to the inflamed jet gen in oxygen. If we breathed in an atmosphere of hydrogen, tould be considered as inflammable, and of course a combustible ngement which I have adopted of classifying as basacigen bodies, ich have heretofore been treated as supporters, with the addition of eas, renders it unnecessary to resort to the incorrect division into re and combustibles.

NITROGEN.

d of distinguishing Degrees of Oxidisement, derived from the School of Lavoisier.

The method which, in concurrence with Thenard, I have purlesignating in the case of the compounds formed by the basacigen rith radicals, the proportion of the former ingredient has been 756.)

In the case of oxacids another method was adopted by the Lavoichool, which, with some modification, still endures, and which I

te as it now prevails.

Agreeably to the nomenclature in question, where, in consequence ent degrees of oxidizement, substances form two acids, one contemporation, the other a lesser proportion of oxygen, the acid, having a proportion, is distinguished by the name of the substance oxy, and a termination in ous; that containing the larger proportion of is designated in the same way, substituting ic for ous; as sulphurland sulphuric acid. That ingredient in an acid or a base, which electro-negative, is called the radical. When an acid is discovered less oxygen than one with the same radical, of which the name the ous, the word hypo is prefixed. Hence the appellations, hypohyposulphurous. The same mean of distinction is employed to a degree of oxygenation exceeding that designated by ous, but a that designated by ic. Hence the name hyposulphuric. If there aid having still more oxygen than the one of which the name ends a letters oxy are prefixed.

. Acids of which the names terminate in ous, have their salts disad by a termination in ite. Acids of which the names end in ic, saits distinguished by a termination in ate. Thus we have and nitrates, sulphites and sulphates. If the base be in excess, deb is prefixed, as subsulphate. If the acid be in excess, super is , as supersulphate. The letters bi are placed before the name of ving a double proportion of acid; hence carbonate and bicarbonate. . The oxide in which the oxidizement is supposed to be at the m is called the peroxide. This monosyllable, per, is also used in of acids, to signify the highest state of oxygenation, and has been thatituted for oxy in the case of perchloric acid. Many chemists apmonosyllable in question to distinguish a salt formed with a peroxhas the red sulphate of iron has been called the persulphate of iron. ate of the red oxide of mercury, the pernitrate of mercury. Agreesimilar rule, salts formed with protoxides have the word proto in the inconces of protonitrate, protosulphate, &c.

has alread been stated that by the British chemists the binary chlorine, bromine, iodine, fluorine, and cyanogen,

et acid, are de ted by the termination in ide.

1058. The word oxide has been erroneously used as a correlative of the word acid, instead of being used as a generic name for any compound oxygen, whether an acid or base. I should deem it preferable to apply the termination in ide, to all compounds of the basacigen bodies, whether acids bases or neutral, employing the words acid and base as terminations to indicate the subordinate electro-negative, and electro-positive compounds. In that case oxybase, chloribase, fluobase, bromibase, iodobase, cyanobase sulphobase, selenibase, telluribase, would stand in opposition to oracis chloracid, bromacid, iodacid, cyanacid, sulphacid, selenacid, telluracid (862, &c.) Yet for convenience, the generic termination ide might be used without any misunderstanding; and so far the prevailing practice might remain unchanged. Resort to either appellation would not, agreeably to custom, be necessary in speaking of salts or other compounds analogous to them; since it is deemed sufficient to mention the radical, as if the sal consisted of an acid combined with a radical, not an oxide. say sulphate of lead, not sulphate of the oxide of lead. This last mentioned expression is resorted to, only where great precision is desirable. cases, it might be better to say sulphate of the oxybase of lead.

1059. The method of indicating the proportion of oxygen in an oxide by changing the termination from ous to ic, has been generally adopted only in the case of the protoxide, and bioxide of nitrogen, the former being usually called nitrous oxide, the latter nitric oxide. In the Berzelian nomenclature, this method of discrimination has been extended to all the compounds formed with amphigen and halogen elements. Hence we have chlorure mercureux, and chlorure mercurique, for the protochloride, and bichloride of mercury; and again, oxide mercureux and oxide mercurique for the protoxide and bioxide of the same metal. These Berzelian names translated into English would make mercurious chloride and mercuric

chloride, mercurious oxide and mercuric oxide.

1060. It should be understood that the employment of the terminations in eux and ique, which in French answer for ic and ous in English, is extended, by Berzelius, to the case of all oxides whether acids or bases. These words are, in my opinion, neither agreeable to the ear, nor sufficiently definite and descriptive. In the received nomenclature, besides the case above cited of the bioxide of nitrogen, the only other instance, of the employment of the letters ic to designate an oxide, is that of the protoxide of carbon called carbonic oxide.

Of the Origin of the erroneous Idea of a Ponderable Acidifying Principle.

was considered as the sole acidifying principle, whence its name as already stated. (637.) Of course, every acid being supposed to consist of oxyget in part, it was enough to call it an acid to convey a correct idea of its composition in that respect. But when, at a subsequent period, it was shown that many acids were destitute of oxygen, and that other substances were nearly as efficient as oxygen in generating acids by a union with acidinal bodies, it became necessary to prefix syllables in order to distinguish the acid compounds produced by one acidifying principle, from those produce by others. (856, &c.) The term acidifying principle originated with the error of assigning that character exclusively to oxygen. From convinience, more than any conviction of its propriety, it was afterwards used o

mally is refered to chlorine, hydrogen, and other elements which found to produce acids by combining with a variety of substances. t he chrious that there is no adeque reason for considering any ileufic element as an acidifying p

Subsequently to the creat of two rd oxygen, the word radithe employed to designate an or the substance. It has since been to all substances with the nigen bedies.

Of Acidity.

1063. Acidity and sourness were originally synonymous. some of the older chemists, the solvent power of ceracid or sour liquids, was ascribed to the sharpness of wer of penetrating and severing the combinations of ber particles was attributed. With people in general, n words acid, and acidity, still retain their original signintion; but by modern chemists, substances are associadas acids which are destitute of sourness, and which extremely discordant in their obvious properties. Thus have in the group of acids, sulphuric acid and flint, negar and the tanning principle; also the volatile and lonferous liquid called prussic acid, and the unctuous, soluble, inert, concrete material for candles, called maruric acid. It might naturally excite the curiosity of the urner, to know by what common characteristic subances so discordant had been affiliated. It would be inmed that there should be some test of acidity, by which determine whether a new compound should belong to e class of acids or not. I am utterly ignorant of any her common characteristic, in these otherwise hetegeneous substances, besides that common preserence the poles, or "electrodes," of the Voltaic series, on ith I have founded my definition of acidity and basid-7; coupled with the inference, mentioned in a note, any compound capable of neutralizing a base, is much to be an acid; and vice versa, any compound public of neutralizing an acid, is deemed to be a base. To me it is quite evident that it is only upon one the other of these characteristics, that many organic strains to be designated as they are.

1664. Amoi 3 the characteristics of acidity heretofore ied on, is that of reddening vegetable blues. By the soluble acids, this property is generally possessed, although an aqueous solution of sulphurous acid is said to whiten litmus, the vegetable blue is generally employed at a test of acidity. But indigo is not reddened by any acid although by nitric acid it is destroyed. Solubility, though usually a property of acids, is in many cases wanting, in those of margaric and stearic acid, and others of similar origin. The acid properties of silicic, and boric, acid are displayed at temperatures incompatible with any other solubility, than that which is effected by the agency of caloric.

Of Alkalinity.

1065. Among the metallic oxides which, agreeably to the definitions above given, are considered as bases, there are a certain number which are called alkalies, on account of some peculiarities which I shall proceed to mention.

1066. All the alkalies have a peculiar taste, called alkaline. They all produce, in certain vegetable colours, characteristic changes, which differ according to the matter subjected to them, but are not varied by changing the alkali.

1067. They restore colours changed by acids, and are

capable of neutralizing acidity.

1068. Acids neutralize alkalies, and restore colours destroyed by them. Acids do not usually combine with acids, nor alkalies with alkalies, but acids and alkalies unite energetically with each other.

1069. By the reaction of alkalies with oils, soaps are

generated, which are soluble in water.

1070. Besides the alkalies above named, there are four other metallic oxides, those of magnesium, barium and strontium for instance, which have been called earths, and which, in different degrees of intensity, have all the alkaline properties abovementioned, excepting that, if not insoluble, they have an inferior solubility, and that they do not form soluble soaps.

1071. There are also some vegetable compounds which possess, to a sufficient extent, the attributes of alkalies, to

be classed among them.

1072. According to Bonsdorf, the halogen elements of Berzelius produce bases, which in some cases display alkalinity. He has noticed a change of colour, indicating an alkaline reaction, on litmus paper, reddened previously by

n acid, and dipped into a solution of a chloride, either of

alcium, magnesium, or zinc.

ranic polarity, are due to some inscrutable influence of the imponderable cause, or causes, of heat, light, and electrity. To a like influence I ascribe the sweetness of sugar, the pungency of mustard or pepper, and of essential oils, well as the endless variety of odour with which these that mentioned products are endowed. It is evident that in the organic alkalies and acids, alkalinity and acidity are found to be associated with combinations of ponderable elementary atoms, which exist in other combinations without inducing alkalinity or acidity.

1074. It is my intention, as introductory to the subject of ammonia, to adduce a few experiments which illustrate

the properties of alkalies in general.

Experimental Illustrations of the characteristic Effects of the Alkalies on certain Vegetable Colours.

1075. Into infusions of turmeric, alkanet, Brazil wood, and rhubarb, a few drops of solutions of either of the alkalies being introduced,—turmeric, from a bright yellow, becomes brown; rhubarb, from nearly the same yellow, becomes red. Brazil wood, from a light red, becomes violet-red; and alkanet, from red, becomes blue. Acids being added, the colours are restored, but by a sufficient quantity of alkali are changed, as in the first instance, and by acids again restored; so that the experiment may be repeated several times with the same infusions.

1076. A blue infusion, obtained from red cabbage, is rendered green by an alkali. By adding some acid, the blue colour is restored; by a further addition of the acid, the infusion becomes red. An alkali being next introduced, it becomes blue, and by a further addition of alkali, the green colour reappears. By alternately using acids and alkalies, these changes may be repeated several times.

1077. The power of various acids in reddening infusions of litmus, shown; and, subsequently, the restoration of the bue colour by an alkali.

COMPOUNDS OF NITROGEN WITH HYDROGEN.

Of Ammonia.

1078. As substances which are analogous in their memoriant properties, are often utterly different in the composition, it is impossible to adopt any arrangement treating of them, which may be in both respects satisfatory. The compound which is the subject of this artic was naturally associated with the other alkalies, whetheir composition was unknown; although now general ranged with the compounds of nitrogen, whilst its form associates are placed among the metallic oxides.

1079. This classification has become the more prop as agreeably to the view latterly presented by Berzeli it appears to be doubtful, whether ammonia be an alka But of this I shall speak more fully, in treating of amn

nium. (1106, &c.)

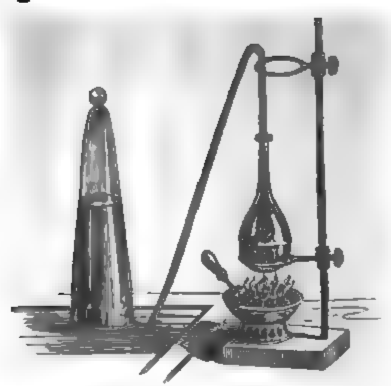
1080. Formerly, besides ammonia, only two other all line substances were known, soda and potash, or potast These being difficult to vaporize, obtained the name fixed alkalies, while ammonia being naturally aëriform, we called the volatile alkali.

- 1081. A new mineral fixed alkali was discovered 1817, and named lithia. It was procured from a stocalled Petalite. Hence its name from the Greek ALL stony.
- 1082. Preparation of Ammonia.—Ammonia is obtain from sal-ammoniac, the salt from which it received name.
- 1083. To evolve this alkali in the gaseous state, equatrized of sal-ammoniac and quicklime, both finely pulvized, are to be heated gradually in a glass matrass. Tammonia is partially extricated by the mere mixture the materials; but heat is necessary to complete the opration.
- 1084. Sal ammoniac, according to the opinion general entertained, is a compound of chlorohydric or muriatic at and ammonia. The lime having a greater affinity for acid than the ammonia, by simple affinity combines we it, and liberates the alkali as a gas, the state which it a turally assumes when isolated. A different view of t subject is taken by Berzelius, which will be explained whereating of ammonium. (1109, 1110.)

85. When it is an object to have the gas perfectly from humidity, it is necessary to arrest the process as as moisture begins to condense in the neck of the rer; or to interpose, between the neck, and the reciused to receive the gas over mercury, a tube conug dry hydrate of potash in small fragments.

'xperimental Illustration of the Process for obtaining Gaseous Ammonia.

96. A flask, containing equal parts of quicklime and mmoniac, both well pulverized and thoroughly interled, is exposed to as much heat as the glass will bear. 67. A bell glass is so placed over the mercurial cisas to receive any gas which may pass from the orist a tube, luted at one end into the flask charged with naterials, and at the other entering the mercury so as under the bell. This apparatus is represented in oflowing cut.



188. Properties of Ammonia.—Ammonia acts like an important properties of taste, upon vegetable colours, in neutralizing acidity. A very small proportion of gas, diffused in the air, is intolerable to the eyes and the of respiration; yet when extremely dilute, the ris agreeably stimulating. Its specific gravity is 18, and 100 cubic inches weigh 18.28 grains. It is inflammable in the air, yet inflames with chlorine teneously, and with oxygen, by the aid of an electric

spark, or galvanic ignition. A candle flame is at first en larged and afterwards extinguished by immersion in this gas. Water absorbs it with surprising velocity, and will hold from 450 to 670 times its bulk. Ice melts in it most speedily than in a fire.

1089. Heat either decomposes, or volatilizes, all ammoniacal compounds; and either of the fixed alkalies, or a the three more powerful alkaline earths, disengage ammonia from any of the acids with which it may be combined

1090. Ammonia, by refrigeration alone, may be condensed into a liquid at — 40° F. By a pressure of six at mospheres and a half, Mr. Faraday succeeded in liquelying it at the temperature of 50° F.

1091. The decomposition and analysis of ammonia have been attempted by ignition with oxygen gas. I have often caused a mixture of it with oxygen, to inflame by means of a wire ignited by galvanism. I believe it to be almost impracticable to ascertain the result accurately by measurement, on account of the liability of ammonia to be absorbed by the moisture of the apparatus, the water produced by the combustion, and the mercury employed to confine the gases.

1092. A spontaneous and explosive c mbustion ensues between chlorine and the hydrogen of gaseous ammo When chlorine is passed in bubbles through concentrated liquid amm ia, a reaction takes place with so much noise, as apparently to endanger the containing vessel. This process has already been mentioned as one of the means of obtaining nitrogen.

1093. In its reaction with ammonia iodine differs from chlorine. When iodine is brought in contact with dry ammoniacal gas, it forms a thick black fluid, which, when saturated with ammonia, becomes more liquid. This compound is decomposed by water forming the iodide of nitrogen. (1032.)

1094. With various metallic oxides, ammonia forms explosive compounds; especially those known as fulminating gold, and the most dangerous species of fulminating silver. By these appellations, however, other compounds of those metals are designated. By some inexplicable influence, probably electro-chemical, the affinities between the oxygen and hydroger are suspended without being destroyed. Yet by slight causes, whether me chanical or chemical, the equilibrium is subverted with explosive violence.

Experimental Illustrations.

1095. Sal-ammoniac and quicklime, being powdered, an mixed in small glasses, pungent fumes are emitted. Am monia extricated by the process above described, and collected in bell glasses over mercury. The introduction cafew drops of water causes the gas to disappear. Icc, i the same way introduced is liquefied, and causes a like result. Characteristic changes effected in the colour of was

er, tinctured by turmeric, alkanet, Brazil wood, and rhu-

- 1096. Evolution of gas shown by means of potash and an ammoniacal salt, introduced into a glass vessel over mercury.
- 1097. Equal volumes of ammonia and chlorohydric cid, mixed, and condensed into a solid, constituting salmoniac.
- 1098. Ammonia inflamed with oxygen gas: also with thorine.
- 1099. Synthesis of ammonia by nitric oxide and hydrogen, heated with platina sponge.

Of the Composition of Ammonia.

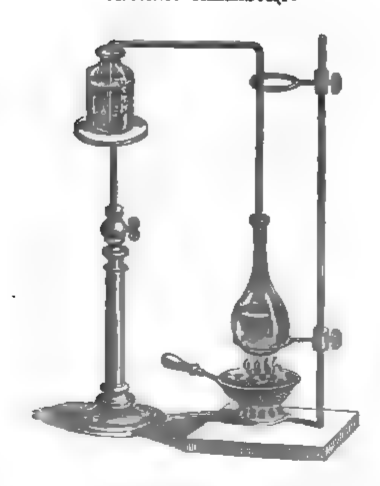
1100. According to Berzelius, ammonia was first ascertained to be a compound of nitrogen and hydrogen, by his celebrated countryman, Scheele. At a later period, Berthollet ascertained the ratio in which these substances that in it, which is by volume that of three of hydrogen to one of nitrogen, condensed into two volumes: and by weight, 3 of hydrogen to 14 of nitro-

gen. See Table, page 189.

- 1101. The partial decomposition of ammonia may be effected by subjecting it to a succession of electrical sparks. Each spark causes the decomposition of a portion of the gas; but as the process proceeds, it becomes more difficult, so that a complete decomposition is impracticable. That portion which is decomposed, is doubled in volume; since the three volumes of hydrogen and one of nitrogen occupy, while combined, but half of the space which they would fill if uncombined.
- 1102. Ammonia, by being made to pass through tubes at a red heat, is resolved into its constituents. This result is promoted by the presence of metallic wire. Any metal will have more or less effect, but iron is most efficacious. It appears from recent experiments of Despretz, that this metal, by continued exposure, may be made to take up nearly twelve per cent. of its weight, becoming a nituret by the absorption of the nitrogen of the ammonia. It is supposed that other metals, which, after a like exposure, exhibit no increase of weight, successively receive and abandon nitrogen; an operation which appears to be singular and mysterious. The metals become brittle during this process. Probably their influence is in its nature electrothemical. In its effects it appears to be the reverse of that by which the mion of the elements of water is promoted by the presence of some metals in a state of minute division.

Process for obtaining Water from Ammonia.

1103. If instead of being conveyed into a bell glass over mercury, the gas be received in water contained in a phial, the water may be saturated, constituting aqua ammoniæ, or water of ammonia. The saturation may be effected in an apparatus, similar to that represented in the following cut.



1104. The absorption of ammoniacal gas by water, causes so much heat, that it is difficult to produce a saturated solution, without assisting the refrigeration by means of ice.

1105. Water saturated with ammonia, when gradually cooled to the temperature of — 40° F., crystallizes in long needles having a silky gloss. No doubt these crystals owe their existence to the presence of water, which exists in them as water of crystallization. Water of ammonia is lighter than water. In combining with the gas, the water loses weight in proportion to the degree of impregnation. At the maximum, at ordinary temperatures, the alkali constitutes about one-third of its weight.

Of Ammonium.

1106. It is well known that Davy resolved potash and soda severally into metals and oxygen, by exposing those alkabes to the divellent influence of the Voltaic current. Subsequently, Berzelias, not having at command an apparatus sufficiently powerful, when unassisted, to effect this decomposition, ascertained that, by placing mercury in contact with a moistened fixed alkali, and in communication with the negative pole, while the alkabicommunicated with the positive pole, an amalgam would result either of potassium or sodium, according to the alkabicinployed.

1107. The results, when ammonia is subjected to the galvanic circuit in contact with mercury at the negative pole, having a perfect analogy, as respects the production of an amalgam, with those obtained by a similar exposure of the other alkahes, as abovementioned, led naturally to the

merence that the causes were analogous; and that, in the case in question, no less than in the others, a metallic radical had been deoxidized and united with the mercury. This inference was rendered more plausible by the evolution of oxygen at the positive pole during the formation of the amalgam. Yet ammonia was known to consist of hydrogen and nitrogen; and to consider either or both of these as oxides, was inconsistent with all the knowledge otherwise obtained respecting them. By some chemists, however, nitrogen was conjectured to be the oxide of a metal, with which this amalgam was supposed to be formed. For this supposed metal, the name of nitricum was suggested. Hence the contact of the amalgam with water was conceived to cause the absorption of oxygen by the nitricum, and consequently the extrication of hydrogen.

1105. Gay-Lussac and Thenard explained the formation of the amalgam, by supposing the absorption of ammonia by the mercury, together with a partion of hydrogen derived from the simultaneous decomposition of water.

1109. Berzelius admits the fact of the union of the elements of ammonia and hydrogen with the mercury, in the proportions alleged by the distinguished philosophers above named; but conceives that, by the addition of an atom of hydrogen to the ammonia, this alkali is converted into a metal, which he calls ammonium. To the union of this metal with mercury, he ascribes the production of the amalgam; and to a resolution of the metal into its elements, the evolution of the ammonia and hydrogen. When an atom of ammonia is presented to an atom of water, he infers that the hydrogen of the water converts it into ammonium, which is simultaneously exidized by the exygen. Hence an atom of ammonia, when combined with an atom of water, may be considered as acting as an oxybase of ammonium. When gaseous ammonia is presented to chlorine, one portion of it is decomposed, of which the nitrogen is liberated, while the hydrogen converts another portion into ammonium. This forms with the chlorine a chloride of ammonium, and, accordingly, by this appellation, sal ammomac, or muriate of ammonia, must be designated, agreeably to the hypothesis under consideration.

1110. When in the process already given for obtaining ammonia, chloride of ammonium (sal ammoniae) is mingled with the oxide of caldim (lime.) by double electrice attraction, the chlorine combines with the calcium, and the oxygen with one atom of the hydrogen in the ammonium; so that water and ammonia are evolved. The latter assumes the gaseous form, while the water unites with the chloride, and remains in union with the heat be not raised unnecessarily, and continued too long.

1111. If we attempt to decompose ammonia without the assistance of mercury, it yields nothing but hydrogen and nitrogen; yet, to produce the analgam, it is sufficient that the wire employed be coated with mercury. The globule of mercury which is left after the spontaneous decomposition which the mass sustains, is in volume surprisingly minute comparatively with the amalgam which it contributed to form.

1112. The most convenient mode of obtaining the ammoniacal amalzum is to place a globule of the amalgam of potassium in a cavity of a prece of chloride of ammonium, slightly moistened. The globule soon enlarges to many times its previous dimensions, by the absorption of the mononium, which relinquishes its chlorine to the potassium.

1113. The ammoniacal amalgam, agitated in dry atmospheric air, yields hydrogen and ammoniacal gas. The same gaseous substances are extriented from it when plunged into ether or naphtha. The ammoniacal amalgam may be preserved for some time, if surrounded by hydrogen, or included in a dry and well closed bottle. When thus protected, and the absence of water is insured by the presence of a small proportion of potas-

sium, it may be kept unchanged for several months.

1114. Berzelius does not consider ammonia as capable of becoming a base, without first being converted into ammonium by the acquisition of hydrogen. In this state, without further change, it can, like other metals, form a salt by combining with any of the halogen substances. But to combine with oxacids, the ammonium must, like other metals, be exidized. The presence of water at once metallizes and oxidizes ammonia. The hydrogen converts the ammonia into a metal, while the oxygen converts that metal into an oxide.

1115. When gaseous ammonia precipitates, from an aqueous solution of a haloid salt,* a metal in the state of oxide, water is decomposed, the hydrogen converting the ammonia into the metal ammonium, while the oxygen converts the metal into an oxide. Meanwhile, the ammonium, combining with the halogen element of the haloid salt, takes the place previously occupied by the metal which has been oxidized.

1116. Agreeably to the view taken above, water, by its contact with ammonia, at once metallizes and oxydizes it, since the hydrogen converts it into ammonium, while the oxygen, at the same time, converts it into an oxide. Thus the formula of ammonia united to water, would be N H^{*} × H^{*};

but when it is resolved into N $H^4 \times 0$, an oxide of ammonium.

1117. It must also follow, that it is not by ammonia that the part of an alkali is performed when entering the arena of alkaline reaction; with the aid of water a transformation takes place, so that the oxide of ammonium is really the ammoniacal alkali. Of course ammonia cannot, consistently

with this explanation, be considered as an alkaline gas.

1118. I deem it expedient to adopt the Berzelian doctrine, as it is necessary to the symmetry of our classification both as respects acids, bases, and chlorides. To consider ammonia, per se, as forming salts with oxacids, or with the halogen bodies, would involve an anomalous deformity, as in all other cases of the union of inorganic acids and bases, the same basacigen ingredient exists both in the acid and the base.

Experimental Illustrations.

- 1119. In a cavity, made in a bit of muriate of ammonia, in communication with one of the poles of the Voltaic pile, a moistened globule of mercury is supported. The mercury is made to communicate with the other pole. The metal swells rapidly, and assumes all the characteristics of an amalgam.
- 1120. An amalgam of potassium, being introduced into a cavity in a piece of sal ammoniac, is rapidly converted into the ammoniacal amalgam, with a prodigious enlargement in bulk.

^{*} A salt formed by a halogen element. (636.)

SECTION III.

OF PHOSPHORUS.

1121. Preparation.—Phosphorus is obtained from the phosphate of soda in urine, or the phosphate of lime in bones. Impure phosphoric acid may be extricated from the earth of bones, by the stronger affinity of sulphuric acid. As, at a high temperature, charcoal takes oxygen from phosphorus, the phosphoric acid is decomposed by ignition with it in a retort, the beak of which is so introduced into water, as to have the orifice a little below the surface. Phosphorus distils into the water, and condenses in tears.

1122. Agreeably to another process, the phosphate of soda, which may be procured at the shops, is decomposed by nitrate of lead, by complex affinity. The phosphorus is separated from the resulting phosphate of lead, by distillation with charcoal, as in the process above men-

tioned.

1123. Properties.—Phosphorus is often of a light flesh colour, but when pure is colourless and translucent. It is rather harder than wax, but is more easily divided by a knife. Phosphorus melts at 108°, and inflames at 148°. At 550° it boils, and may be purified by distillation from a retort filled with hydrogen gas, receiving the product under water. Phosphorus is insipid and probably inodorous; but, in consequence of its oxidizement, it emits a feeble alliaceous odour of phosphorus, or hypophosphoric acid. When pure it is flexible, but the presence of 1-600th of sulphur renders it brittle. Its specific gravity is 1.77. Subjected to the rays of the sun, it acquires a red colour. If heated to 155° and suddenly cooled, it becomes black. Thenard, however, states that this change cannot be effected in phosphorus which has not been repeatedly distilled. He suggests it as possible, that the colour of phosphorus, when pure, is black; and that the colour which it usually assumes, may be due to the presence of hydrogen, which has been long known to be evolved, when phosphorus, in the usual state, is fused and subjected to the Voltaic current.

1124. Exposed to the air at ordinary temperatures, phosphorus combines slowly with oxygen, appearing luminous in the dark, but without any sensible evolution of heat. Less heat is requisite to cause the inflammation of phosphorus

phorus in atmospheric air than in oxygen; and less als is necessary in this last mentioned gas, in proportion a the pressure is reduced. When sprinkled with powdere sulphur, carbon, fluoride of calcium, carbonate of lime and various other bodies, and placed in a receiver from which the air is subsequently exhausted, phosphorus in flames. Professor Alexander D. Bache, who has muce enlarged the list of substances capable of producin this result, has succeeded in inflaming phosphorus in a exhausted receiver by enveloping it in muslin, or in pape pierced with small holes. He conceives that, with the exception of bodies exercising a chemical affinity, as in the instance of sulphur, the substances associated with the phosphorus act mechanically, and have upon it no other effect than that of promoting its union with the oxygen remaining in the receiver. This opinion is corroborated by the fact that the removal of the air may be too rapid, or too complete, to produce the inflammation.

in boiling naphtha, by gradual refrigeration. Like sulphur, phosphorus, in volatilizing, produces a feeble light, without entering into any chemical combination. Water in which phosphorus has been kept, oxygen being excluded, acquire the power of shining when agitated. The admission of air destroys this phosphorescent property. Phosphorus is oxidized by the action of nitric or nitrosonitric acid, and

converted into phosphoric acid.

Experimental Illustrations of the Properties of Phosphorus

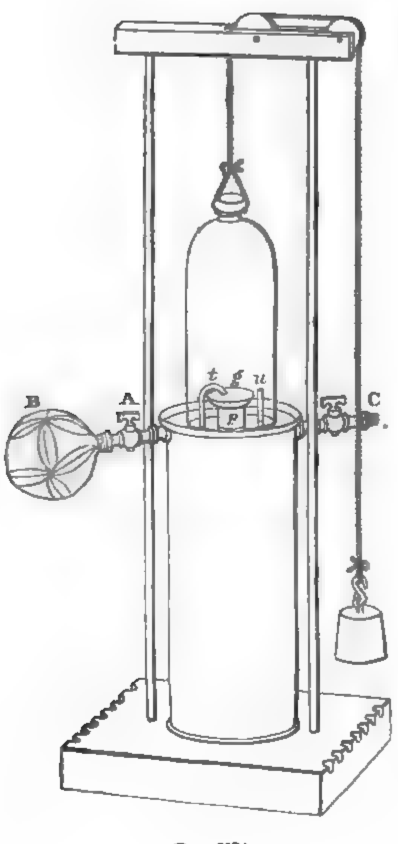
a gentle heat. Luminous appearance in the dark. Combustion in oxygen, (654,) in nitrous, and nitric oxide, unde hot water by a jet of oxygen, and by nitrosonitric acid (1131.)

1127. Anomalous combustion of phosphorus consequento rarefaction.

Combustion of Phosphorus in Nitric Oxide.

1128. The backwardness of the gaseous oxides of nitrogen to part with their oxigen to substances, under circumstances in which it would be readily yielded by a mospheric air, has been already mentioned, and a method of illustrating it has bee described. (969.) The opposite engraving represents an apparatus, which may lused to extend the illustration to nitric oxide, which, producing a corrosive fume nitrous acid by admixture with oxygen, cannot be employed in apparatus requiri the aid of an air-pump, without corroding the metal of which such instruments a partially constituted. The apparatus in question is nearly the same as that used to

Combustion of Phosphorus in Nitric Oxide.



(Page 212.)



separation of nitrogen from atmospheric air. There are, however, in this, two issued tables; and the bell comployed is without any cap or cook. The cook at which a gam electic bag, supplied with exygen gas, is attached, communicates a pips, which descends close along the inner lateral surface of the cylindrical or vessel till it reaches the bottom, then bends at right angles, and proceeds the bottom of the vessel till it reaches the copper pipe in the axis of the vessel, it bends at right angles upwards, and ascends vertically in close contact with size, till it reaches the copper cup, g, by which the pipe is surmounted. It is so recurved as to overhang and direct its orifice, t, downwards, into the cavity a copper cup.

9. Another tube, u, proceeds from its junction with a screw and cook, C, on ther side of the vessel, and descends to the bottom, rising again, like the tube smentioned, along the central pipe, till it reaches somewhat above the brim of up, where it terminates without a curvature. After the proper quantity of shorus has been placed in the cup, the atmospheric air may be allowed to enfrom the bell glass through the cook, C, by sinking it into the water, with a the vessel must have been filled nearly to the brim. The air being expelled, communication made with a celf-regulating reservoir of nitric oxide, by means s flexible leaden tube attached to the cook for that purpose, the bell may be sup-with a quantity of this gas, sufficient to occupy about two-thirds of its especity, took being then closed, and the communication with the reservoir interrupted, hot iron must be introduced through the bore of the central pipe, p, till it see the cup. For this purpose, it is of course necessary that the apparatus d be upon a table with a suitable sperture, and of a height sufficient to allow

to outer the orifice of the pipe, p.

Although by the heat of the incandescent iron, the phosphorus will be, no combustion will ensue, until, by opening a communication with the gume bag, a small quantity of oxygen is allowed to enter. But no sooner is this steed to take place, than a most brilliant and almost explosive evolution of heat ight sounce. A higher temperature is requisite to ignite phosphorus in nitric in mittees oxide.

in nitreus czide.

Inaction of Phosphorus with Mitrees-mitric Acid.



1131. If into a tall tube of about an inch and a half in diameter, and fifteen inches in height, some strong nitric acid be introduced, and about five grains of phosphorus, a reaction will curue, which is invariably energetic, and sometimes explosive. The phosphorus abstracting oxygen, the acid is converted into nitric oxide gas and nitrous acid vapour, which are conjously evolved, so as to fill the upper part of the tube, and overflow it with a beautiful red fume. Meanwhile, vivid flashes arise from the oxygenation of the phosphorus, and pieces of it are occasionally thrown up to the case in the tube, where a wind comup into the gas in the tube, where a vivid combustion ensues between the phosphorus, and the exygen of the nitric exide gas or nitrous acid. 1132. The residual nitric acid will be found in-

termingled with phosphoric acid.

1133. Latterly, in performing this experiment, I have surrounded the tube with a very stout glass cylinder, and another of wire gauze; as upon one occasion a violent explosion took place, which did much damage to my apparatus. If the phosphorus be reduced into small fragments, the risk of an explosion is increased. Heeting the acid, before the addition of the phosphorus, ensures an explosive reaction.

Application of Phosphorus to Endomatry.

191. One of the most simple modes of accortaining the quantity of oxygen in the introduce into a graduated tube, standing over water, and containing 100 tures of sir, a stick of phosphorus, supported by a wire. The phosphorus slowly from in the nitrogen, and, combining with the oxygen, condenses with it, and

cause a corresponding absorption of the water. When, by these means, t gen is all removed, the quantity of nitrogen remaining will be known by ins the graduation. The difference between this quantity and 100, the number sures taken, is the quantity of oxygen present.

A Simple Atmospheric Endometer by Phosphorus.



1135. If a cylinder of phosphorus be supports a wire (as represented in the adjoining cut.) w glass matrass, inverted in a jar of water, the of the included air will be gradually absorbe order to determine the quantity of oxygen in two have only to ascertain the ratio of the quantity of air inch the matrace at the commencement of the proce

1136. This object may be attained by weigh matrase when full of water, and when contains portion only which rises into it in consequence absorption. As the weight in the first case is weight in the last, deducting the weight of th in both cases, so will 100 be to the number of 100 of atmospheric air, which consist of cays
1137. Again, the contents of the vessel may

covered by the sliding-rod gas measure, (936,) absorption measured by introducing from the s strument, as much air as will compensate it. whole content to the quantity which compensate sheorption, so is 100 to the quantity of oxygen parts of the atmosphere.

1138. If the neck of a vessel of this kind he one-fourth as much as the bulb,—by gradue neck, so that each division will represent a her

part of the whole capacity, the result may be known by inspection.

1139. Eudiometrical processes by the slow combustion of phosphorus are t requiring many days to complete them, and consequently the aid of baremetr servations to ascertain and allow for any intervening changes in atmospheri

1140. It is alleged that nitrogen is enlarged one-fortieth of its bulk, by the phorus which it dissolves. This is to be deducted in estimating the recision | 1141. The action of the phosphorus may be accelerated by heat; but in the operation must be performed over mercury, and the manipulation will be troublesome and precarious.

1142. I have never in this way, obtained results comparable in accuracy & formity, to those procured by the hydro-oxygen eudiometer. (940, &c.)

Volumescope for the Analysis of Atmospheric Air by Phospharus.

1143. A volumescope has been described, (816.) for showing the diminu bulk in five volumes of atmospheric air, consequent to the admixture of nitric The same apparatus may, with some modification, be employed to show the a tion of volume resulting from the combustion of phosphorus. This object is a by associating with the volumescope, the apparatus employed for the comba phosphorus in oxygen. (654) For this purpose, the volumescope, instead o situated over the pheumatic cistern, should be placed in a small tub, into the of which is inserted a tube, supporting, at the upper extremity, the cup for the phorus. The phosphorus being placed in the cup, and water in the tub, this is raised by an air pump, until no more than five volumes of air remain in the der. The phosphorus is then ignited by means of a red-hot iron, and the conducted as already described. (922.) As soon as the expansion resulting fi heat of the combustion ceases, it will be seen that a little more than one volu of the five has been condensed.

COMPOUNDS OF PHOSPHORUS WITH OXYGEN.

1144. These compounds are four in number; one c oxide of phosphorus, and three acids, hypophosphorous,

phorous, and phosphoric acid. Their composition is as follows:—

equivalent 48, / lent 56.

Three atoms of (with one atom of oxygen, equivalent phosphorus, { 8, form oxide of phosphorus, equiva-

phosphorus, equivalent 32, with one atom of oxygen, equivalent 8, form hypophosphorous acid, equivalent 40.

Two atoms of with three atoms of oxygen, equivalent 24, form phosphorous acid, equivalent 56.

with five atoms of oxygen, equivalent 40, form phosphoric acid, equivalent

Of Oxide of Phosphorus.

1145. When phosphorus, melted under hot water, is subjected to a jet of oxygen from a tube with a capillary orifice, oxide of phosphorus and phosphoric acid are produced. The acid dissolves, and the oxide, being at first suspended in the water, subsides subsequently in red flakes. oxide is insipid and inodorous. It is not luminous in the dark, even when rubbed. At a heat a little below redness in close vessels, it is decomposed into phosphoric acid and phosphorus. If the air be admitted, phosphoric acid is the sole product. The oxide of phosphorus takes fire spontaneously in chlorine, producing the perchloride of phosphorus and phosphoric acid. It is inflamed by the action of nitric acid. With chlorate of potash it explodes violently; also with nitrate of potash previously warmed. white matter with which phosphorus becomes coated when kept in water, and which is generally supposed to be a hydrate of the oxide, is stated by Thenard to be a hydrate of phosphorus.

Production of Oxide of Phosphorus experimentally illustrated.

1146. Production of oxide of phosphorus, by the reaction of oxygen with that substance, while in fusion under hot water.

Of Hypophosphorous Acid.

1147. This acid is obtained by precipitating the baryta from an aqueous solution of hypophosphate of that base. The acid remaining in solution, may be so concentrated by evaporation as to become a vivid liquid, highly acid, and even crystallizable. It is an energetic deoxidizing agent, and forms numerous salts, all of which are soluble in water, whereas several of the phosphates are insoluble.

Of Phosphorous Acid.

1148. This acid has been generally considered as the product of the slow combustion of phosphorus with atmospheric oxygen; but Thenard alleges that this product is a peculiar acid, intermediate in its degree of oxidation between phosphorous and phosphoric acid, and to which he has given the name of hypophosphoric acid. Phosphorous acid may be procured by passing vaporized phosphorus over corrosive sublimate heated in a tube. Chloride of phosphorus results, which, by reaction with water, produces chlorohydric and phosphorous acids. The chlorohydric acid, being more volatile, may be expelled by heat.

1149. Phosphorous acid is a colourless, inodorous, crystalline substance, possessing a pungent taste, and reddening litmus paper. Like hypophos-

phorous acid, it possesses powerful deoxidizing properties.

Of Phosphoric Acid.

by adding sulphuric acid to phosphate of baryta suspended in water. The sulphuric acid unites with the baryta, forming an insoluble salt, which precipitates while the phosphoric acid remains in solution. When phosphorus is gradually added to nitric acid, phosphoric acid is generated, and remains mingled with the residual nitric acid.

rated, and remains mingled with the residual nitric acid.

1151. Properties.—Phosphoric acid is an inodorous, colourless, viscid liquid, possessing in a high degree the property of reddening litmus. It cannot be obtained in a state of liquidity free from water. When exposed to a red heat and afterwards cooled, it forms a transparent brittle glass. This fusion should be effected in a platinum crucible; since phosphoric acid, when heated to redness, attacks either glass or porcelain. The acid, if examined after this exposure to heat, is found, although its composition remains the same, to have acquired new properties. On this account, the name of paraphosphoric has been given to it; while the term phosphoric is applied to designate the acid in the state first described. Nitrate of silver yields with phosphoric acid a yellow precipitate; with paraphosphoric acid a white one. Albumen is coagulated by the latter, but not by the former.

1152. Solid paraphosphoric acid, when exposed to the air, deliquesces, and is in a few days converted into phosphoric acid. The same change is produced in a short time by boiling water. The solid white flakes which are obtained during the quick combustion of phosphorus with oxygen, consist of paraphosphoric acid. It may likewise be produced by fusing the biphosphate of soda, which by these means is converted into a paraphosphate. Mr. Graham, who has made a number of interesting experiments on this subject, states that the acid which is con-

thined in of is a third species of phosphoric n composition with the others, but no this species he has given

the name of pyrophospho.

containing the same n atoms of t suments, and having the life int, t ter meric has been applied. I us, no a and pyrophosphoric acide to

Of the Chlorides of Phosphorus.

in chlorine. If the chlorine be in excess, the perchloride is formed; if the phosphorus be in excess, the sesquichloride is obtained. The sesquichloride is a transparent, colourless, fuming, inflammable liquid, heavier than water, and having a disagreeable smell. When brought into contact with water, a reciprocal decomposit in takes place, and chlorohydric and phosphorus acid are produced. The perchloride is a white, crystalline, inflammable body, which is converted into vapour at a temperature much below 214°. It forms a neutral compound with ammonia, and its vapour is alleged to redden dry litm paper. Hence, by some chemists, it is considered as an acid. I dout whether litmus paper is ever reddened by an acid, unaided by water. The perchloride and water decompose each other, forming phosphorus in these chlorides, as the oxygen bears to the phosphorus in phosphorous, an phosphoric acid.

Of the Bromides and Iodides of Phosphorus.

1155. The sesquibromide is a yellow furning liquid; the perbromide, a crystalline volatile solid. In their reaction with water and composition, they agree with the chlorides of phosphorus. Iodine appears to combine with phosphorus in almost every proportion. There are, however, at least two definite combinations, which correspond in composition with the chlorides and bromides.

Of the Sulphides and Selenides of Phosphorus, commonly called Sulphurets and Seleniurets.

1156. When phosphorus is melted with sulphur, or when sprinkled with it, and placed in a receiver from which the air is subsequently withdrawn, (1124,) a sulphide of phosphorus is formed. This sulphide may consist of various proportions of its ingredients, according to the circumstances under which it is produced. Sometimes it is liquid, sometimes solid.

1157. Selenium, like sulphur, combines with phosphorus in almost every proportion. The sulphides and selenides of phosphorus are decomposed

by water.

1158. The incorporation of sulphur with phosphorus, when effected by heat, is sometimes productive of explosion; and the resulting mass is spontaneously inflammable in the air; being the sole active ingredient in some friction matches.

COMPOUNDS OF PHOSPHORUS WITH HYDROGEN.

Of Protophosphuretted Hydrogen.

1159. Protophosphuretted hydrogen may be obtained by heating a contrated solution of phosphorous acid, or by adding phosphorus to the

torials for generating hydrogen.

1160. Properties.—It is a colourless, inflammable gas, with an osimilar to that produced by the combustion of arsenic. Under the ordinarement of the atmosphere, protophosphuretted hydrogen does not inferenced with oxygen; but, if the pressure be reduced about one-time combustion ensues.

1161. On meeting with oxygen, this gas becomes luminous in the cin consequence of the slow combustion of the phosphorus; though the evolved is inadequate to inflame the hydrogen. If the process for ducing the philosophical candle, (806,) be repeated with the additionsome comminuted phosphorus to the materials, protophosphuretted hydra will be generated, and, escaping into the air, will produce a jet lumino the dark.

Of Perphosphuretted Hydrogen.

chlorohydric acid with the phosphuret of calcium, which is obtained by jecting lime to the vapour of phosphorus at a bright red heat in a parc or coated glass tube. The gas may also be evolved by heating in a re 75 grains of phosphorus, 1500 of slacked lime, with 4 ounces of water 50 grains of caustic potash, and 40 of phosphorus, moistened by 60 c of water. The phosphorus should be atided first, and the potash last the heat which it evolves, contributes to the heat required for the opers. The body of the retort should be filled with hydrogen, or a few drog ether should be added, to prevent the first portions of the gas from influ with the atmospheric oxygen of the retort. By its affinity for the phosphorus, and the metal of the phosphuret, the oxygen of the water is separ from the hydrogen, which, while pascent, unites with a portion of the phorus, and forms perphosphuretted hydrogen.

1163. The following cut represents the apparatus usually employed

obtaining perphosphuretted hydrogen.



1164. The beak of the retort being depressed below the surface of

mercury, each bubble as it escapes into the atmosphere, explodes. It produces at the same time a dazzling flash, which is transformed into a beautiful wreath of smoke, consisting of aqueous vapour and phosphoric acid, created by the oxygenation of the hydrogen and phosphorus. Each wreath, as it rises, expands in diameter, and, when the bubbles succeed each other quickly, a series of them may be seen in the air at the same time.

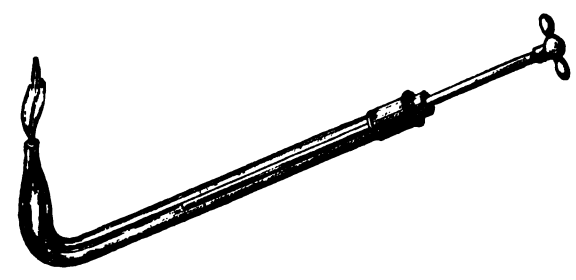
1165. Properties.—Perphosphuretted hydrogen is a colourless gas, possessing an alliaceous smell, and a bitter taste. Water dissolves it in small quantity, forming a yellow solution, which has a bitter taste, and a smell resembling that of the gas. When this gas is brought in contact with oxygen, or atmospheric air, it explodes with a loud noise and a vivid flash; being converted into phosphoric acid and water. The same mixture, in more tubes, undergoes a similar change slowly, and without the evolution of heat and light.

1166. Perphosphuretted hydrogen may be decomposed either by heat, by the electric spark, or by the rays of the sun. Professor Rose considers protophosphuretted and perphosphuretted hydrogen as isomeric, and of course similar in composition, though different in properties. If the opinions of Rose are correct, one should be called phosphuretted hydrogen, the

other paraphosphuretted hydrogen. (1153.)

1167. Chemists do not agree in their statements respecting the composition of the compounds of hydrogen with phosphorus.





1163. This figure illustrates an advantageous employment of the sliding-rod eudiometer, in exhibiting the spontaneous combustion of phosphuretted hydrogen, the splendid colour of the flame of cyanogen, and other experiments, where the combus-

tible character of a small quantity of gas is to be shown.

1169. For the experiments in question, the instrument is charged, agreeably to the mode already described in the case of the eudiometers, by introducing the apex into any bell glass or other vessel holding the gas, and drawing out the rod; by which a portion of the gas, equivalent in bulk to the part of the rod withdrawn, enters the receiver of the eudiometer through the hole in the apex. The receiver being then removed from the bell glass, and held up in a position favourable for observation, the rod is slowly returned into its tube, so as to expel the gas in a jet suitable for inflammation. In the case of perphosphuretted hydrogen, the gas burns spontaneously as soon is it escapes from the apex. In the case of other inflammable gases, inflammation is produced by the flame of a taper.

SECTION IV.

OF CARBON.

1170. Nature presents us with the most beautiful and purest specimens of this substance. The diamond is pure carbon. When equal weights of charcoal and diamond are severally exposed to the focus of a powerful lens in oxygen gas, included in different bell glasses, they are both converted into carbonic acid, from which, by ignition with potassium, carbon may be precipitated.

1171. Carbon is very abundant in nature, in the various kinds of fossil coal, from anthracite or plumbago, in which it is nearly pure, to the variety called candle, or cannot coal, which abounds with bitumen. In bituminous coal there is much hydrogen. Carbon pervades vegetable and animal matter as an essential element. It is, especially, a

constituent of the fibres of wood.

1172. Until of late, plumbago was considered as a chemical compound of iron with carbon. Berzelius alleges it to be carbon mingled, but not combined, with iron and

other impurities.

1173. I ascertained that anthracite, when completely burned in oxygen gas, produced no diminution of volume, the products being water and carbonic acid. I infer, therefore, that the combustible portion of this coal consists almost solely of carbon, united with hydrogen and oxygen in the proportion for forming water. It may, in fact, be deemed an hydrate of carbon.

1174. Preparation.—In the laboratory, charcoal is obtained, sufficiently pure, by heating wood intensely in close vessels. In the large way, it is procured by igniting large quantities of wood, so covered with earth, that the access of air may be at first controlled and afterwards

prevented.

1175. Coke is obtained from bituminous coal, by a process analogous to that employed for obtaining vegetable charcoal, which it resembles in chemical, though not in

mechanical properties.

1176. Properties.—Carbon is inodorous, insipid, and usually black. Charcoal of wood is one of the best radiators, and worst conductors of heat. There is reason for believing this peculiarity to result from its excessive poro-

carbon. 221

etter, and probably radiates it worse. Charcoal is highly

susceptible of galvanic ignition.

1177. Next to the metals, charcoal is the best conductor of electricity. It appears, from the experiments of Proessor Silliman, that charcoal, when exposed to the influence of a powerful Voltaic series, is volatilized, so as to be transferred from the positive to the negative pole, on

which it forms a projection.

1178. Charcoal, when intensely ignited without access of air, becomes denser, harder, and a better conductor of heat. Substituting animal products for those of vegetation, in the usual process of carbonization, animal charmal is obtained. It does not, like the coal of vegetable substances, retain the form of the bodies from which it may be procured, and is replete with cavities, created by the escape of the gaseous elements associated with it in the organic state. It has a grayish-black colour, and a prilliancy resembling that of plumbago. Carbon is precipitated in various forms from coal gas; among others, in that of long brittle filaments, associated in tufts, resembling locks of hair.

diamond, or in that of common charcoal, when examined in the pulverulent form, so that the result shall not be affected by the numerous cavities existing in it when in mass, is about 3.5. The apparent lightness of charcoal is caused by its porosity. The specific gravity of anthracite does not exceed 1.6; that of plumbago is 2.32; yet they are both much more compact than charcoal, and, in proportion to the space occupied by them in mass, ob-

viously much heavier.

1180. Carbon, under some circumstances, appears to have a transcendent affinity for oxygen. In its ordinary state it requires a temperature above redness, in order to exhibit this affinity—in other words, to burn. In proportion as it becomes denser, we find it more difficult to ignite; in proportion as it may be more minutely divided, or approaches a state of extreme porosity, it is rendered nore susceptible of ignition. Thus the susceptibility of gnition increases from the diamond to tinder in the following order:—Diamond, plumbago, anthracite, coke, charoal of hard wood, charcoal of soft wood, tinder. In some

forms, and when mixed with iron, as when obtained by carbonizing Prussian blue, or tanno gullate of iron, it takes

fire spontaneously at ordinary temperatures.

1181. According to Despretz, carbon, during its combustion, evolves sufficient caloric to melt one hundred and five times its weight of ice. It is not to be inferred that this is true of carbon in all its forms. Berzelius alleges that the same degree of heating power is not possessed by every kind of charcoal; some of its forms, according to him, producing much more heat in burning than others. This I should not believe without conclusive evidence.

Of the Decolorizing and Disinfecting Power of Charcol.

1182. Carbon, as procured from organic products, especially animal matter, displays a great power to combine with and precipitate colouring matters. Hence it is attensively used in the refining of sugar, and generally in chemical processes, in which the objects of research are entangled with colouring matter. This power is not inherent in the colouring matter. rent in elementary carbon, but appears to be due to its previous associations, or to some peculiarity of arrangement, derived from the process of carbonization.

1183. Animal charcoal is much more efficacious than that derived from vegetables. The carbonaceous mass, obtained by igniting blood with carbonate of potash, appears to have the greatest efficacy. That the presence of an alkali during the ignition contributes to the effect, seems to justify the conjecture that cyanogen, the generation of which, in combination with the alkali, is a necessary concomitant, has some agency in the process. Charcoal is a powerful antiseptic, operating efficiently in preserving water or meat from putridity. Moreover, water rendered extremely foul, as that from the public sewers, may be purified by filtration through pulverized charcoal. In fact, filters are now extensively manufactured, in which charcoal is the most efficient and only chemical agent employed. The gravel, sand, and sponge, usually associated with it, act mechanically.

Of the Power of Charcoal and other Substances to absorb Aëriform Fluids.

1184. Charcoal, which has, in a state of ignition, been submerged in mercury, on being introduced into gaseous substances, condenses into its carbon. 223

The quantity of the surrounding aëriform matter, whatever it may The quantity condensed varies with the gas, from 90 times the bulk ne charcoal, as in the case of ammonia, to 1.75 times its bulk, as in the of hydrogen. During their absorption, the gases give out heat, and more in proportion to the rapidity with which the condensation is effected if, on the other hand, by exposure within an exhausted receiver, pas be evolved, cold is produced. Charcoal, thus deprived of gas, rerbs any gas exposed to it, as greedily as if recently ignited.

185. This faculty of absorbing gaseous substances, is impaired by huty, which charcoal is prone to absorb in the form of vapour, afterwards lensing it into the state of water. Water partially displaces the gases

iously absorbed.

186. The aëriform fluids, absorbed by charcoal, are expelled by heat langed, with the exception of sulphuretted hydrogen and oxygen. The let deposites sulphur, and the latter is gradually converted into carbonic

The absorption of this last mentioned principle continues for some, but, in quantity, has not been found to exceed 14 times the volume of carbon. In a rarefied medium, charcoal absorbs less in weight, but e in volume; so that the increased resistance of the gas, arising from a nution of pressure, counteracts, to a certain extent, the power of the to condense into its pores a certain weight. The power of absorption es in a degree with the number and minuteness of the pores existing in charcoal; of course, it varies with the wood by which it is yielded. I recoal of box-wood is pre-eminent in absorbing power; that furnished woods of a lighter kind is very inferior in this power. Plumbago and aracite have no capacity, even after ignition, to absorb gases.

1187. In the property of absorbing aëriform fluids, charcoal is not sinar. De Saussure ascertained that different porous minerals, and many ds of wood, also silken and woollen stuffs, absorb many times their vo-

re of gas.

es, they are absorbed in proportion to their reciprocal attractions, and t exercised by the pores of the substances employed. A mixture of igen with hydrogen or carbonic acid, is more copiously absorbed than er when alone; yet by heat or exhaustion they are liberated without ainution. Nevertheless, sulphuretted hydrogen and oxygen, when acted in by charcoal, produce water, sulphur being deposited.

1189. The absorption of moisture by charcoal and other porous bodies long been noticed. On this account, it is difficult to weigh such bodies thout an increase of their weight, even when they are placed in the scale like red-hot. Those aëriform fluids are absorbed to the greatest extent, ich are capable of assuming the liquid state. These facts explain the greatation in weight received by charcoal exposed to the air, which

counts to between ten and twenty per cent-

1190. I have devoted more space to this subject, because it illustrates a openty which otherwise might not be sufficiently considered. It forms a cubar instance of mechanico-chemical agency, if I may be allowed to a new word to express the idea. Without the porous or cellular structe which it possesses in the form of charcoal, carbon is not endowed the either disinfecting, absorbing, or colour-removing powers; and yet is evident that the carbon acts in charcoal by a species of chemical afixy, unaided by which the cellular structure would be inefficient in the remove under consideration. As respects the transmission of contagious

or infectious effluvia, the absorbing power of porous bodies merits a I believe that the carbonaceous matter, evolved during the burning actually neutralizes those fetid emanations which it is employed to in the chambers of the sick.

COMPOUNDS OF CARBON WITH OXYGEN.

One atom or one volume of carbon, equivalent 6,

with one atom or half a volume of o equivalent 8, forms one atom of volume of carbonic oxide, equivalent with two atoms or one volume of o equivalent 16, forms one atom of volume of carbonic acid, equivalent

1191. Two atoms or volumes of carbon, equivalent with three atoms or one and a half volumes of o equivalent 24, form one atom of oxalic acid, equivalent

1192. Two other compounds of carbon with oxyg alleged to exist; one called *mellitic*, the other *croconi* The former contains four atoms of carbon to three of gen; the latter five of carbon to four of oxygen.

Of Carbonic Oxide.

1193. Preparation.—This compound is produced combustion of carbon with an inadequate supply o gen; or when bodies containing carbonic acid are with certain substances having an affinity for or Thus it may be procured by heating carbonate of with iron filings. The best process, however, for or ing carbonic oxide in a state of purity, is to heat five of concentrated sulphuric acid with one of oxalic which, being deprived by the sulphuric acid of the which is essential to its existence, is resolved into bonic oxide and carbonic acid. The latter gas me removed by lime-water, leaving the carbonic oxide state of purity.

Apparetus for separating Carbonic Acid from Carbonic Oxide, by means of Lie

1194. This apparatus is represented by the opposite engraving. Lime-wa introduced in sufficient quantity into the inverted bell glass, another smallass, C, is supported within it as represented in the engraving. Both of have perforated necks. The inverted bell is furnished with a brass cap stuffing box attached to it, through which the tube, D, of copper, slides About the lower end of this tube, the neck of a gum elastic bag is tied; so cavity of the bag may communicate with that of the tube. The neck of bell is furnished with a cap and cock, surmounted by a gallows screw, by which the leaden pipe, P P, with a brass knob at the end suitably perform be fastened to it, or removed at any moment. Suppose this pipe, by aid of brass knob at the other extremity, to be attached to the perforated neck tall bell glass filled with water upon a shelf of the pneumatic cistern: on

Apparatus for separating Carbonic Acid from Carbonic Oxide, (Mouns of Lime-roater.



(Page 224.)









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nication between the bells, the water will subside in the tall bell glass over a, and the air of the bell glass, C, being drawn into it, the lime-water will and partially occupy the space within the latter. As soon as this is efse cocks must be closed, and the tall bell glass replaced by a small one 1 water, and furnished with a gallows screw and cock. This bell being atthe knob of the lead pipe, to which the tall bell had been fastened before, atus is ready for use. I have employed it in the new process for obtaining oxide from oxalic acid, by digestion with sulphuric acid in a glass retort. ous product consists of equal volumes of carbonic oxide and carbonic acid, ing received into a bell glass, communicating, as above described, by a pipe bell glass, C, may be transferred into the latter, through the ripe, by openocks. As the gaseous mixture enters the bell, C, the lime-water subsides. s a sufficient quantity of the gas has entered, the gaseous mixture, by means m elastic bag and the hand, may be subjected to repeated jets of lime-water, depurated of all the carbonic acid. By raising the liquid in the outer bell, rified carbonic oxide may be propelled through the cock and lead pipe, into el to which it may be desirable to have it transferred.

5. Properties of Carbonic Oxide.—Carbonic oxide is urless, insipid gas, indecomposable by heat or electronic, and incapable of reddening litmus. Its specific is 0.9727. It does not support combustion, and is ctive to life. It burns with a feeble blue flame, and, ning with oxygen, is converted into carbonic acidatinum sponge, a mixture of oxygen and carbonic is gradually changed into carbonic acid.

Experimental Illustrations.

6. Carbonic oxide gas, evolved from oxalic acid by ocess abovementioned, and collected in bell glasses vater. Combustion and detonation of it with oxyis, effected by means of a sliding-rod eudiometer, or escope. Subsequent absorption of the resulting caracid by lime-water, shown.

Of Carbonic Acid.

- 7. The proportion of this gas, existing in the atmosis much less than was formerly supposed; being, acgreg to some experiments of Thenard, not more than a undth part. It is this portion, however, that prothe pellicle on lime-water, during its exposure to the nd which, under like circumstances, by combining the alkalies, enables them to effervesce with acids. nic acid is incessantly a product of combustion and respiration of animals. It is a principal ingredient rble and limestone.
- 18. Preparation.—Carbonic acid may be evolved from 29

any carbonate by heat or by acids. It is use lly procured for the impregnation of water, by the superior affinity of sulphuric acid for the lime in marble. Excepting that it is more costly, chlorohydric acid is preferable for this purpose; as the chloride of calcium, being very soluble, does not, like the sulphate, clog the vessels.

1199. Carbonic acid is evolved copiously during the vi-

nous fermentation.

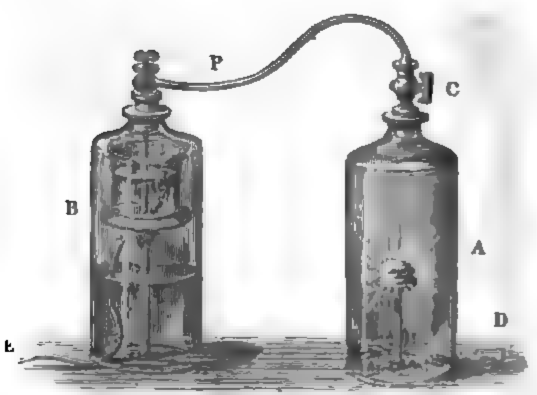
1200. The process and the self-regulating reservoirs, already described, (796, &c.) may be resorted to for carbonic acid, substituting lumps of marble for zinc. The best materials for the evolution of this gas, agreeably to my experience, are chlorohydric acid and calcareous stalactites, or clam shells.

1201. Carbonic acid might be procured at a triffing cost, by drawing, by the aid of a suction pump, the efficient of burning charcoal through water to deprive it of dust, and then forcing it into the cavities in which its pre-

sence may be desirable.

1202. This process for the production and employment of carbonic acid, generated by the combustion of charcoal, is illustrated in the small way by the following engraving and description.

Combustion of Charcoal or other Combustibles in Ozygen Gas.



1263. The preceding cut represents an apparatus which I have contrived for a hibiting the combustion of charcoal, or other combustibles, in oxygen gas. To large glass bells, A, B, each furnished with a tubulure at the apex, are associated by

carbon. 227

ns of the pipe, P, which, in one of the bells, B, communicates with a tube, exing about five inches within the bell, below its neck, so as to reach into some -water, er an infusion of litmus, contained in a glass vessel, resting on a stand, epresented in the figure. The wooden stand which holds the glass vessel, and iron stand which supports the coal in the bell, A, must be previously placed on shelf of the pneumatic cistern, as represented in the cut; so that A, when ining the coal, may be over the mouth of the cock, D, which communicates with a the gas holders, situated under the shelves of the pneumatic cistern, which,

his experiment, should be filled with oxygen.

M. Into the bell glass in which the vessel is placed, a pipe from the suction pump e hydrostatic blowpipe is made to enter, and reach nearly to the stand. The ratus having been prepared thus far, the bells must be lifted so as to permit a soal to be put upon the iron stand, as represented in the figure. As soon as are restored to their previous situations, the suction pump must be put into ation, and the cock, D, of the gas holder, containing the oxygen, opened; so as ow a current of the gas to have access to the coal, by replacing the air, which thdrawn by the pump through the pipes, P and E. The coal burns splendidly; as the oxygen becomes saturated, it is drawn off by the suction pump, being in its way from A to B, to pass through the liquid in the vessel, into which ends the tabe proceeding from A. If the liquid be water tinged with litmus, it become red by the action of the carbonic acid: if it be lime-water, a copious y precipitate will appear.

205. Properties of Carbonic Acid.—It is a colourless, with a pungent smell and an acid taste. Water takes its own bulk of this gas, whatever may be its density. It nbines with earths, alkalies, and metallic oxides, formwith lime, baryta, strontia, magnesia, and oxide of d, compounds which are insoluble. Hence it precipies lime-water, barytic-water, and solution of acetate of d. Litmus is reddened by this acid. It destroys life d extinguishes flame, but is not insalubrious to breathe len much diluted with air.

1206. Carbonic acid is very antiseptic. When concented in water it is grateful to the stomach. Potassium rns in this gas, absorbing oxygen and precipitating carn. Plants probably absorb it, retain its carbon, and re out its oxygen. The respiration of animals tends compensate this change, by carbonizing the oxygen of

e air.

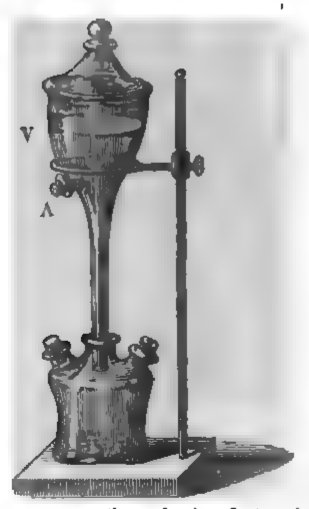
1207. Carbonic acid is heavier than atmospheric air, specific gravity being 1.5239. At the temperature of ?, and under a pressure of forty atmospheres, it connects into a colourless liquid.

Experimental Illustrations.

1208. Evolution of the gas shown; also its property of stinguishing a candle. That it differs from nitrogen, ade evident by means of lime-water. Litmus, reddened carbonated water, and restored to its original colour boiling.

1209. Analysis of mixtures containing the ga means of the sliding-rod eudiometer and lime-water.

Apparatus for showing some of the distinguishing Properties of Carbonic Sci



1910. Having introduced in three-necked bottle, represented adjoining figure, one or two or carbonate of ammonia, add she as much nitroso-nitric acid. (101 active ofervescence will enme, from the expulsion of the carbo from the ammonia, by the stree finity of the nitrie acid. At the time, sufficient fume will be to make it evident how far the are occupied by the gas, to the sion of atmospheric air. By thes the movements of the carbonic s will be recognised as ascending apper vessel, which it will it finally overflow through the cre-tween the brim and cover.

1211. The cover being runn lighted candle will couse to burn lowered into the fume indicati space occupied by the gas. The will comprise the whole cavity vessel, so long as the sporture closed; but, on removing the estimate aperture, the gas will flow the streets marked by the streets. the streem, marked by the acce ing fume, will be seen descend wards the table, and will extingulation of a candle if made to en it; or, it may be received into

so as to arrest the combustion of a taper introduced into it, or upon which tents of the mug may be poured. Under these circumstances, a taper will be where within the vessel, V, if it be not below the aperture, A, above which is not now seen to extend itself. But if one of the orifices of the bottle be c the carbonic acid will be found entirely to desert the upper vessel,

1212. It will thus be made evident that this gas, from its greater specific; has, in the atmosphere, some of the habitudes of liquids; while its incape support combustion will be demonstrated.

1213. The specific gravity of carbonic acid being rather more than a

greater than that of atmospheric air, it does not speedily leave any cavity if may be introduced. It is on this account that persons often perish on a wells.

Impregnation of Water with Carbonic Acid.

1214. The process by which water is impregnated with carbonic acid,

easily understood from the following engraving
1215. A condenser, A, is fastened at bottom into a block of brass, which nished with a conical brass screw, by means of which it is easily attached f the floor. In this brass block are cavities for the two valves, one opening the floor. It the other outwards towards the pipe, C. The pipe, B, nicates with a self-regulating reservoir of carbonic acid,

1216. The gas which the condenser draws in from the reservoir, is forced the other pipe into a strong copper vessel, in which the water is situated, an is represented in the figure, as if the front part were removed, in order to cal-

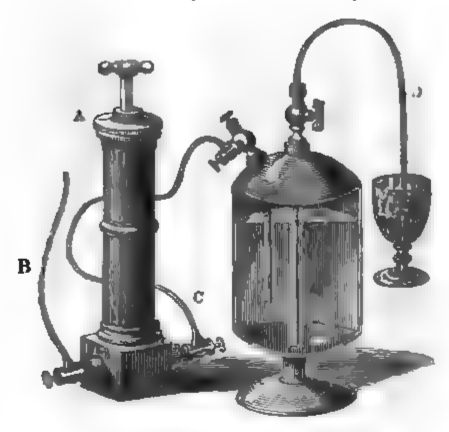
inside to inspection.

1217 If the vessel and its contents be thoroughly exhausted of air before pregnation is commenced, the water will take up as many times its bulk of the pressure employed exceeds that of the atmosphere.

1218. When duly saturated, the water may be withdrawn at pleasure by a

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the syphon, D, of which one log descends from the vertex of the vessel to the bottom, while the other is conveniently situated for filling a goblet.



Of the Liquefaction and Solidification of Carbonic Acid.

1219. It has been shown that the extrication of carbonic acid from a base, may be checked by the pressure consequent to confinement, (242,) and it has been mentioned that Faraday obtained this acid in a liquid state, by causing the materials for the generation of it to react within a glass tube, sealed hermetically after their introduction. Subsequently, the liquefaction of this acid was accomplished on a much larger scale by Brunel; and in 1836, thirteen years after the date of Faraday's observations, Thillorier caused not only the liquefaction, but the solidification of the acid. Without any other knowledge than that afforded by brief notice, or verbal information conveyed by travellers, my friend, Dr. Mitchell, was quite successful in the repetition of the processes of Thillorier. The production of the solid acid is dependent on the same principles as the congelation of water in the cryophorus and in Leslie's experiment. (309, &c.)

1220. The pressure requisite to retain carbonic acid in a state of liquidity, is at 4° below zero, 26 atmospheres; at 32°, 36 atmospheres; at 86°, 75 atmospheres. (196.) Its specific gravity is, at that temperature, about 830. The density of the gas which occupies the cavity above the liquid portion of the acid, is 130 times the density of that which it has at the mean baro-

metric pressure of 30 mches of mercury.

1221. Liquid carbonic acid does not combine, nor even mingle, with water or fixed oils; but, under the requisite pressure, combines readily with ether, alcohol, naphtha, or oil of turpentine. It may be decomposed by

potasium, but not by zinc, iron, or other metals proper-

1222. One of the most interesting properties of the acid, is that intense cold produced by its assuming the aëriform state, to which allusion has been made. A jet of it depressed a thermometer to 130° below zero, F. The cold by which the acid is frozen, or in other words, its freezing point, is estimated at 148° below zero, F. According to Mitchell, one drachm of

solid acid is yielded by each ounce of the liquid. I will here give Dr. Mitchell's description of this wonderful product of chemical art, in his eva words:*

1223. "The porosity and volatile character of the solid renders its specific gravity of difficult ascertainment. When recently formed it is about the weight of carbonate of magnesia, and w in strongly compressed by the fingers, its density is nearly doubled. Soud carbonic acid is of a perfect whiteness, and of a soft and spongy textue, very like slightly moistened and aggregated snow. It evaporates rap y, becoming thereby colder and colder; but the coldness produced seer to steadily lessen the evaporation, so that the mass may be kept for some time. A quantity weighing 346 grains lost from 3 to 4 grains per minute at first, but did not entirely disappear for 3 hours and a half. The natural temperature was 76° - 79°. The solid is most easily kept when compresed and rolled up in cotton or wool. Its temperature when newly for a is not exactly ascertainable because it is immediately lowered by eval ation. Thillorier seems to have entertained the opinion that the gr is gree of cold was created at the time of the formation of the solid. In my experiments a constant decrease of temperature was observed; which was accelerated by a current of air, or any other means of augmenting evaporation. At its formation, the carbonic snow depresses the thermometer to about — 85°. If it be confined in wool or raw cotton, its cooling influence is retarded; if it be exposed to the air, especially when in motion, the thermometer descends much more rapidly; and under the receiver of an air pump, the effect is at its maximum. The greatest cold produced by the solid carbonic acid in the air was - 109°, under an exhausted receiver - 136°, the natural temperature being at $+86^{\circ}$.

1224. "The admixture of sulphuric ether so as to produce the appearance of wet snow, increased the coldness, for the temperature then fell, under exhaustion, to —146,* a degree of cold which we were not able to exceed by means of any variation of the experiment. That result is most easily obtained by putting about two fluid drachms of other into the iron receiver before charging it. A compound liquid may be thus formed which yields a snow in less quantity, but of a more facile refrigeration. Alcohol may replace ether in either mode, but with less decided effect. In the air the alcoholic mixture fell to —106° and remained stationary. By blowing the breath on it, it fell to —110°. Left to itself it rose slowly to —106°; but

on being placed under an exhausted receiver sell to -134°.

1225. "Every attempt to wet the carbonic solid with water, failed, so that

no estimate of its relative effects could be made.

1226. "The experiments resulting from the great coldness of the new solid, were very striking. Mercury placed in a cavity in it, and covered up with the same substance, was frozen in a few seconds. But the solidification of the mercury was almost instantly produced by pouring it into a paste made by the addition of a little ether. Frozen mercury is like lead, soft and easily cut. It is ductile, malleable, and insonorous. Just as it is about to melt, it becomes brittle or 'short' and breaks under the point of a knife. These facts may account for the discrepancies of authors on this subject. Frozen mercury sinks readily in liquid mercury.

1227. "At about — 110° liquid sulphurous acid is frozen, and the ice

^{*} For engraving and description of Mitchell's modification of Thillorier's appearatus, see Appendix.

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anks in its own liquid, and at — 130° alcohol of .798, assumes a viscid and oily appearance, which by increase of cold, is augmented until at — 146° it is like melted wax. Alcohol of .820 froze readily. At — 146°

rulphuric ether is not in the slightest degree altered.

1228. "When a piece of solid carbonic acid is pressed against a living nimal surface, it drives off the circulating fluids and produces a ghastly thite spot. If held for 15 seconds it raises a blister, and if the application e continued for two minutes a deep white depression with an elevated marin is perceived; the part is killed, and a slough is in time the consequence. have thus produced both blisters and sloughs, by means nearly as prompt s fire, but much less alarming to my patients."

Of Oxalic Acid.

1229. Latterly oxalic acid, long known as a product of regetation, has been found to belong to the compounds of carbon with oxygen; and still more lately mellitic and croconic acid have been added to this class. Yet when the necessity of water to the existence of these acids is taken into view, it appears to me questionable whether they may not be considered as acids with a compound radical, con-

sisting of hydrogen and carbon.

1230. Preparation.—Oxalic acid may be obtained from the common sorrel, Rumex acetosa, or from the wood sorrel, Oxalis acetosella, from which it derives its name. In these plants it exists in the state of binoxalate of potash. It may also be procured by the reaction of one part of sugar with six of nitric acid. The weight of the acid obtained is equal to three-eighths of the materials. Wood, glue, silk, or hair may be substituted for sugar in this process; but when these substances are used, the product is impure. Next to sugar, starch and molasses are probably the best materials. Oxalic acid may be procured also, by digesting shavings of wood in a solution of caustic potash, at a heat considerably above that of boiling water.

1231. Properties.—Oxalic acid is a solid, but soluble both in water and alcohol, the resulting solutions being extremely sour. One grain in half a pint of water is sufficient to redden litmus distinctly. It cannot exist uncombined with water or some other base. The atomic composition of this acid would authorize us to consider it as a binary compound of carbonic acid and carbonic oxide. In every atom of oxalic acid in its appropriate crystalline form, there are three atoms of water. When these crystals are exposed to an unusually dry atmosphere, or to a temperature of 80°, a partial efflorescence ensues; and if the heat

be raised to 212°, they part with two atoms of wat which they recover on exposure to the air after cooling

When heated to 300°, the acid is decomposed.

1232. Oxalic acid is an energetic poison. The best tidotes for it are magnesia, or the calcareous carbona in the pulverulent form, especially chalk. acid meets with either of these bases, an insoluble a inert oxalate is formed. Hence its employment as a t

for lime.

1233. It appears from statements made by Vogel in t Journale de Pharmacie, for April, 1836, that the protoxid of iron and copper are precipitated from their union wi sulphuric acid by oxalic acid. The oxalate of iron is y low; the oxalate of copper, blue. Both are insoluble water.

Of Mellitic Acid.

1234. Mellitic acid is obtained in crystals from a re mineral, called the honey-stone, which is a mellitate alumina. It is soluble in water and alcohol, and has sour taste.

Of Croconic Acid.

1235. Croconic acid may be procured in yellow crysta from the croconate of potash, which is generated in the pr cess for obtaining potassium by means of charcoal. inodorous, has an acid and astringent taste, and redde litmus.

COMPOUNDS OF CARBON WITH OXYGEN AND CHLORINE.

1236. There are two compounds of carbon with oxygen and chlori To one of these, which has been recently discovered, the name of chlo has been given; to the other, that of *chlorocarbonic* or *chloroxycarbo* acid. The latter name is preferable; as the other would convey the i of an acid made solely by the union of chlorine with carbon.

Of Chloral.

1237. When chlorine is passed through alcohol, which consists of drogen, oxygen, and carbon, one portion combines with hydrogen, form chlorohydric acid, while another combines with oxygen and carbon, so ing chloral.

1238. Chloral is described as a colourless transparent liquid with a p gent odour. Its specific gravity is 1.502. It boils at 201°, and may distilled unchanged. With water it forms a white crystalline mass, a

rently a hydrate.

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Chloral consists of nine atoms of carbon, four of oxygen, and six

Of Chloroxycarbonic Acid.

When one volume of dry chlorine and one volume of carbonic are mingled, and exposed to the solar rays, they combine, and into one volume of a colourless acid gas, to which the name of arbonic acid has been given. It is exceedingly offensive to the to the organs of respiration. It reddens litmus paper, and with forms a white salt. By contact with water a reciprocal decompuses, and chlorohydric and carbonic acids are produced. It conce atom of chlorine, and one atom of carbonic oxide.

Of the Chlorides of Carbon.

Thlorine forms four compounds with carbon. The dichloride is ystalline inflammable solid, having a peculiar odour, resembling ermaceti. At 250° it sublimes in crystals. It is fusible by heat, at a temperature between 350° and 450°. The dichloride cone atom of chlorine and two of carbon.

When the liquid, produced by the union of chlorine with olefiant I bichlorine ether, is exposed to the sun, in contact with a sufficitity of chlorine, the sesquichloride of carbon is produced. It is so, transparent, friable, crystalline body, nearly tasteless, and recamphor in smell. While exposed to the flame of a spirit lamp, with a red flame, but the combustion ceases as soon as the lamp d. It melts at 320°, and at 360° is converted into vapour, which ondensed without decomposition. It is nearly twice as heavy as the sesquichloride of carbon consists of three atoms of chlorine toms of carbon.

The protochloride is obtained by passing the sesquichloride in rough a red-hot porcelain tube. The sesquichloride is decomthe protochloride and chlorine. The protochloride is a transparless liquid, with a specific gravity of 1.4875. It is composed of of chlorine and one of carbon.

All the above described chlorides are insoluble in water, acids, and out are soluble in oils, alcohol, and ether. When chloral is boiled on of potash, a decomposition ensues, and a chloride of carbon in vapour, and may be condensed in a receiver. This chloride less, transparent liquid, with an odour similar to that of chloric specific gravity is 1.48. This chloride consists of five atoms of ad four of carbon.

Of Bromide of Carbon.

When bromine is brought in contact with half its weight of perarbon, heat is evolved, a decomposition ensues, and bromides of carbon are formed. The bromide of carbon is a volatile, colour, of a sweet taste, and an ethereal odour.

Of the Iodides of Carbon.

The protiodide of carbon is a liquid, in properties strongly rethe bromide of carbon. The periodide appears under the form

of yellow crystalline scales, which have a sweet taste, a strong arountic smell resembling that of saffron, and a specific gravity higher than that of water.

Of Sulphocarbonic Acid, or Bisulphide of Carbon.

1247. The bisulphide of carbon is obtained by passing the vapour of sulphur over charcoal heated to incandescence in a porcelain tube. It is a transparent, colourless, volatile liquid, possessing an acrid taste, and a peculiar nauseous smell. Its specific gravity is 1.272. It boils at 105°, and does not freeze at — 60°. At a temperature a little above the boiling point of mercury, it inflames. When the bulb of a spirit thermometer, wapped in lint imbued with this liquid, is placed within a receiver, and the air withdrawn, the temperature falls to —82°.

1248. This compound unites with almost all the sulpho-bases, forming with them sulpho-salts, and is as well entitled to be treated as an acid, as

the analogous compound formed by sulphur with hydrogen.

1249. Dr. Thomson supposes that the solid mass, obtained by washing the nitre out of gunpowder, is probably a solid sulphide of carbon.

COMPOUNDS OF CARBON WITH HYDROGEN.

1250. Carbon and hydrogen are in opposite extremes, as respects their susceptibility of the aëriform state. Per se, carbon is probably more difficult of volatilization by heat, than any other substance in nature. Hydrogen, on the other hand, as far as our experience goes, is not susceptible of condensation, even into the non-elastic state of fluidity. There is, however, a powerful affinity between these substances; and hence, when a compound which contains them is subjected to heat, they are made to combine in various proportions, according to the intensity of the ignition, and the influence exercised by the nitrogen, or oxygen, previously in combination with them.

1251. In general, the compounds of carbon with hydrogen are distinguished by inflammability. In the gaseous state they constitute, when ignited, the flame of candles, lamps, gas lights, and culinary fires. They are incapable of supporting life, but are not actively noxious when di-

luted with the air.

1252. The gaseous compounds of carbon with hydrogen are obtained by the destructive distillation of bituminous coal, wood, oil, tar, and other inflammable substances.

1253. The illuminating power of each of these various kinds of gas, seems to be in proportion to the quantity of carbon contained in a given volume, provided there be an equivalent supply of oxygen; but, otherwise, the excess of carbon renders the flame smoky. Hence the greater bril-

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liancy of small flames, or those excited by a current of air, as in the Argand lamp. The same flame which in common air is unpleasantly fuliginous, transferred to oxygen gas, displays a perfect brilliancy.

1254. The known compounds of carbon with hydrogen are numerous and complicated; and yet it is probable that many exist in nature, or may be produced by art, with

which we are at present unacquainted.

1255. We have had occasion to state, (1153,) that where bodies have, in the same volume, the same number of atoms of each of their ingredients, and yet differ in their properties, they are said to be isomeric, from ires equal, purpos part. Compounds, in which the constituents are in the same ratio, but in which the resulting volumes exist in different degrees of condensation, are said to be polymeric with respect to each other, from reduction, purpos part. The last term is applied to a class of the compounds of carbon with hydrogen, in all of which these elements exist in the same ratio of atom for atom; yet from some difference in the mode of aggregation, or, as I believe, in the extent and modes of their association with heat, light, and electricity, their degree of condensation when in the aëriform state, and their properties in other respects are quite different.

1256. We have then two groups of the carburets of hydrogen, in one of which diversity of properties is attended by a corresponding diversity in the ratio of the carbon to the hydrogen; while in the other this ratio is uniform, although the properties and resulting volumes in the aëriform state,

differ. In the first group, there are four compounds.

1257.—1. Light carburetted hydrogen, or fire damp, consisting of two volumes or atoms of hydrogen, with one volume or atom of carbon.

1255.—2. The compound, in all the varieties of which there are as many atoms of one element as of the other, and for which Dr. Thomson proposes the name of carbohydrogen as a generic appellation.

1259.—3. Bycarburet of hydrogen, in which six atoms of carbon are

united with three of hydrogen.

1260.—4. Naphthaline, in which ten atoms of carbon are combined

with four atoms of hydrogen.

1261. The second group, which is subordinate to the first, being formed in fact by the ramifications of carbohydrogen, comprises, according to Dr. Thomson, several varieties, which he designates and describes as follows:—

1262.—1st. Protocarbohydrogen, consisting of a volume of carbon and a volume of hydrogen, condensed into one volume. This variety, now called mytheline, has been lately isolated by Dumas and Peligot, by distilling one part of pyroxylic spirit, obtained by the distillation of wood, with two parts of chlorohydric acid, and three of sulphuric acid; when an ethereal chlorohydrate of mytheline results. Subjected to a red heat, this ethereal compound is resolved into chlorohydric acid gas, and mytheline in the gaseous form. Pyroxylic spirit is considered as a bihydrate of mytheline, being procured from crude pyroligneous acid by distillation. It bears the same relation to mytheline that alcohol does to etherine. (1267.)

1263.—2d. Deutocarbohydrogen, or olefiant gas, consisting of two vo-

lumes of carbon and two of hydrogen, condensed into one volume.

1264.—3d. Tritocarbohydrogen, consisting of three volumes of carbon

and three of hydrogen, condensed into one volume. This is by Dr. Themson considered as constituting the gas evolved from oil, which was by Dalton called super-olefiant gas.

1265.—4th. Tetartocarbohydrogen, consisting of four volumes of car-

bon, and four volumes of hydrogen, condensed into one volume.

1266.—5th. Hexacarbokydrogen, containing, according to Thomson,

six volumes of each element, condensed into one volume.

1267. Of Etherine.—Besides these compounds, it has been inferred, by many chemists, that there is a liquid, or solid compound, formed of four volumes or atoms of carbon, and four volumes or atoms of hydrogen, condensed into one volume or atom. This has been called etherine, under the idea that it is the common base of all the ethers, forming common ether by uniting with one volume of aqueous vapour, alcohol, by uniting with two such volumes, and the various ethers, by uniting with acids, or the other ingredients, after which they are severally named. Etherine would of course be isomeric with the tetartocarbohydrogen of Dr. Thomson. (1265.)

Of Light Carburetted Hydrogen, or Fire Damp.

1268. The substance distinguished by these names has been dignified by a variety of appellations, among which are heavy inflammable air, carburetted hydrogen, and bihydroguret of carbon. Dr. Thomson has, is some instances, used the monosyllable di to indicate proportions the inverse of those indicated by the monosyllable bi. Thus, bichloride of carbon would signify two atoms of chlorine and one of carbon, while dichloride conveys the idea of two atoms of carbon and one of chlorine. Consistently, then, I think, Dr. Thomson should have called this gas, a dicarburet of kydregen; as the proportions of its constituents are the inverse of those in the bicarba ret. This gas has long been known to miners of bituminous coal, under the name of fire damp, as one of their greatest enemies. It is liberated co piously from cavities in the coal, in which, no doubt, in many instances, it has been pent for ages. It is also evolved from the mud of stagnant waters. and is occasionally emitted from fissures in the earth. There is no good mode of forming it artificially. It is a colourless gas, of course irrespirable but having more than a negative influence in destroying life. Its specific gravity is 0.5593.

Of the Safety Lamp.

1269. In the account above given of dicarburet of hydrogen, it was mentioned that it was in mines a source of injury. When existing in the air beyond a certain proportion, it explodes on coming into contact with the flame of a lamp or candle Hence, as artificial light is necessary in mines inaccessible to the light of day, the use of candles or lamps, in the ordinary way, has been frequently destructive to the workmen. It had, of course, been the cause of great misery to them, and of em

barrassment to the proprietors of the mines.

"steel mill" had been resorted to, in which the rapid revolution of a steel wheel agains a flint, was made to produce a succession of sparks, and of course a feeble light, believe that the security afforded by this invention was imperfect, and the light is sufficient. Explosions have been more frequent in the English mines of late year probably in consequence of the greater extent and depth to which they are excavate. While under the painful impression made by some recent catastrophes of this nature in which many miners had been been killed or mutilated, Sir H. Davy exerted his self to discover the means of sustaining flame safely within explosive gaseous mittures. He soon ascertained that his object might be effected by enclosing the flat in a cage of wire gauze, so as to allow of no communication with the surroundit medium, which does not take place through the meshes of the gauze. Owing to t

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ing power of the wire, the mixture cannot pass through the meshes in a state of bustion. Of course the inflammation is confined within the wire game.

71. The method in which I illustrate the operation of the safety lamp, may be y comprehended from the following figure. The lamp is seen within a large



glass cylinder upon a stool. The cylinder is closely covered by a lid, which will not permit the passage of air between it and the cylinder, and which is so light as to be easily blown off. Excepting the cage alluded to above, the safety lamp does not differ materially from those which are ordinarily used. The upper surface of the receptacle for the oil, forms the bottom of the cage, which is so closely fitted to it, and so well closed every where, as to allow air to have access to the flame only through the meshes of the wire gause. The cage is enclosed within three iron rods, surmounted by a cap, to which a ring for holding the lamp is attached, as seen in the figure.

1272. If, while the lamp is burning,

1272. If, while the lamp is burning, as represented in the figure, hydrogen, either pure or carburetted, he allowed, by means of the pipe, to enter the glass cylinder, so as to form with the air in it an explosive mixture, there will nevertheless be he explosion. It will be found that es the quantity of inflammable gas increases, the flame of the lamp enlarges, until it reaches the wire gauze; where it burns more or less actively, accordingly as the supply of atmospheric air is greater or less. It will, under these circumstances, often appear as if the combustion had ceased; but on increasing the

rtion of atmospheric air, the flame will gradually contract, and finally settle the wick, which will burn as at first when the supply of hydrogen ceases.

73. If the cage be removed from the lamp, and the experiment repeated in all respects as at first, an explosion will ensue, as soon as a sufficient quantity of ogen is allowed to enter the cylinder.

Deutocarbohydrogen, or Olefiant Gas, called also Carburetted Hydrogen, and Hydroguret of Carbon.

1274. This gas received its name in consequence of its ag condensed with chlorine into a liquid, having an aginous consistency, although otherwise unlike an oil. was discovered in the year 1796. It may be obtained subjecting a mixture of five parts of sulphuric acid with e of alcohol to heat in a glass retort. It is invisible, d possesses, like other gases, the mechanical properties atmospheric air. Its specific gravity is 0.9808. When awn into the lungs it produces asphyxia. It burns with metaplendour, and detonates with oxygen with such vioce, that without some precautions it is dangerous to

analyze it by the usual processes. I have had several eudiometers broken by it, but have latterly avoided that accident, by exploding the mixture in a rarefied state, into which it is easily brought in some of the instruments

which I employ.

1275. The analysis may be performed in the volumescope for analyzing the air by means of hydrogen, with a degree of accuracy sufficient for the purpose of illustration. Four volumes of oxygen should be added to one of the gas. The ignition being effected as already described in the case of pure hydrogen, it will be seen that the five volumes are reduced to less than three, and that by the introduction of lime-water, these three may be reduced to one residual volume of oxygen. The reason why the residual gas is less than three volumes, is, that the carbonic acid formed is partially absorbed by the water. gas contains in one volume, two volumes of hydrogen, and two of carbon vapour, it will, for the latter, require two volumes; for the former, one volume of oxygen. Of course the hydrogen, and the oxygen which combine with it, will be condensed; so that after the explosion, unless so far as absorbed by the water, two volumes of carbonic acid will remain mingled with the one volume of oxygen in excess.

Of certain Gaseous Compounds formed by igniting the Gascous Elements of Water, while containing Olefiant Gas, or the Vapour of Ethers, or Essential Oils.

1276. I observed some years ago, that when olefiant gas is inflamed with an inadequate supply of oxygen, carbon is deposited, so copiously as to render the glass receiver of the eudiometer impervious to light, while the resulting gas occupies double the space of the mixture before explosion. Of this I conceive I have discovered the explanation. By a great number of experiments, performed with the aid of my barometer-gauge eudiometer, I have ascertained that if during the explosion of the gaseous elements of water any gaseous or volatile inflammable matter be present, instead of condensing there will be a permanent gas formed by the union of the nascent water with the inflammable matter. Thus two volumes of oxygen, with four of hydrogen, and one of olefiant gas, give six volumes of permanent gas, which burns and smells like light carburetted hydrogen. The same quantity of the pure hydrogen and oxygen, with half a volume of hydric ether, gives on the average, the same residue. One volume of the new hyponitrous ether, under like circumstances, produced five volumes of gas.

1277. An analogous product is obtained when the same aqueous clements are inflamed in the presence of an essential oil. With oil of turper-tine a gas was obtained, weighing, per hundred cubic inches, $16\frac{3}{10}$ grs., which is nearly the gravity of light carburetted hydrogen. The gas obtained from olefiant gas, or from other, weighed on the average, per the

carbon. 239

me bulk, $13\frac{5}{10}$ grs. The olefant gas which I used, weighed per hunred cubic inches, only $30\frac{5}{10}$ grs. Of course, if, per se, expanded into six olumes, it could have weighed only one-sixth of that weight, or little over we grains per hundred cubic inches. There can, therefore, be no doubt not the gas obtained by the means in question is chiefly constituted of ater, or of its elements, in the proportion in which they exist in that liquid. we table, page 189, for steam.

1278. The gas created in either of the modes abovementioned does not nation carbonic acid, and when generated from olefiant gas, appears by nalysis to yield the same quantity of carbon and hydrogen as that gas

fords before expansion.

1279. These facts point out a source of error in experiments, for anayzing gaseous mixtures by ignition with oxygen or hydrogen, in which be consequent condensation is appealed to as a basis for an estimate. It prears that the resulting water may form new products with certain vola-

ilizable substances which may be present.

1280. The gas obtained by passing the vapour of alcohol through an gnited porcelain tube, is confounded generally with that which results from he reaction of sulphuric acid with alcohol, as above described, (1273,) but equal volumes of the gaseous products obtained, the two processes being analyzed, I found that procured by ignition to have only condensed half as nuch oxygen as the other. From the facts above stated, that the presence of water causes a union between its elements, and those of the carbon and hydrogen of carburets, whether in the form of vapour or gas, it may be inferred that the products of the decomposition of alcohol must vary accordingly as it may be more or less anhydrous. The alcohol which I employed was of the specific gravity nearly of 840°: were absolute alcohol subjected to the process in question, a gas containing a larger proportion of carbon might be obtained. (619, 1252.)

Experimental Illustrations.

1281. Cork, cotton-seed, caoutchouc, and nuts, introduced in small quantities into a gun-barrel, of which the butt-end has been heated to a bright red-heat. Brilliant jet of flame proceeds from the touch-hole. Inflammation of the gas extricated by distillation from oil or bituminous coal, also of olefiant gas. Olefiant gas, mixed with oxygen gas, and exploded in a sliding-rod cudiometer. Residue renders lime-water milky.

Of Gas Lighting.

122. The gaseous compounds of carbon and hydrogen have been much applied to the purpose of illumination.

12:3. The gas, for this purpose, is obtained by the destructive distillation of bituminous ead, oil, or resinous substances, and is received in gasometers, whence it is distributed through pipes to the burners. (617, 1252.)



1984. One of the gradient obstaces to the grand employment of gas lights as a an situte for cash and lamps, is the necessity of pipes leading free gas ometers to all situations where the light is wanted The condensation of the gas into strong metallic re-ceivers, has been resorted to in order to obviate the difficulty. This process may be illustrated by mean of the apparatus described for the impregnation of water with carbonic acid, being medified as sup-

ed in the adjoining cut.

1285. It is only necessary to exchange the summa-nication with the self-regulating reservoir of automic acid gas, for a similar communication with a men ar of elefant gas; and the copper vessel being fat ex-hausted of air, to condense the gas into it. The sy-phon used for the affect of the impregnated water is replaced by a cock and tube, the latter terminaling in a capillary perforation. Through this, the gumay be allowed to escape in a proper quantity to protect a gas light when inflamed. It has, however, shows appeared to me, that the expense of condensing the gas, and of procuring and transporting the reserve would render this method of affording light disable.

1996. Latterly, the loss of gaseous matter, by es-

process unprofitable.

Of some Varieties of Carbohydrogen, and of the Bicarburet of Hybr-

1287. Tetartocarbohydrogen, hexacarbohydrogen, and bicarburet of bydrogen were all obtained by Mr. Faraday from the liquid which is deposited from oil gas, when condensed into vessels under great pressure for the purposes of illumination.

1288. On subjecting the matter, deposited as above described, to a very gentle heat, tetartocarbokydrogen is separated in the form of a transparent, colourless, inflammable gas, with a specific gravity of 1.9444. When cooled to zero, it condenses into a transparent colourless liquid of the specific gat-

vity of 0.627, being the lightest liquid known.

1289. When the liquid remaining after the extrication of the tetarious bohydrogen is heated, vapour is evolved, and the boiling point continually rises until the temperature of 176° is attained. Between this temperature and 190°, a large portion distils in the form of a liquid. When this liquid is cooled to zero, it separates into two compounds, one of which becomes The liquid is the compound when solid, while the other continues liquid. Dr. Thomson calls hexacarbohydrogen, though its composition does not ap pear to have been well ascertained. It is inflammable, soluble in alcoholand boils at 176°. The solid compound is the bicarburet of hydrogenis at ordinary temperatures a colourless, transparent, volatile liquid, which boils at 1862, and has a specific gravity of 0.85. At 322 it crystallines, and, when cooled to zero, acquires a consistency like that of loaf sugar.

Of Naphthaline.

1290. Naphthaline is obtained by subjecting to distillation the tar which is formed during the decomposition of bituminous coal. The first products are ammonia water, and the liquid called coal naphtha; but towards the close of the process, naphthaline is obtained.

241

290. Naphthal white crystalline substance, with an aromatic

and a pungeou assegreeable taste.

.291. There are other compounds of carbon and hydrogen,—native shths for instance, and oil of turpentine. The almost endless variety of essential oils derived from vegetables, consist either wholly or principal-scarbon and hydrogen. Of some of these I shall hereafter briefly treat; stice them all would be inconsistent with the limits prescribed to this k.

If the Compounds formed by (bon with Chlorine and Hydrogen.

292. It has already been stat d that olefant gas received its name onsequence of its being conde sible with chlorine into a liquid of an ginous consistency. To this liquid the name of chloric ether has been roperly given, as it indicates a dependency on chloric acid for its constituence of generation, contrary to the fact. As it consists of two atoms of prine and one of etherine, a more appropriate name would be bicklorine er.

1293. Bichlorine ether is limpid and colourless like water, has a pleasant

ell, and an agreeable sweet taste.

294. Chlorine combines with several other of the polymeric varieties of bohydrogen, forming with them compounds of different properties. It produces two compounds by combining with the bicarburet of hydro; one solid, the other liquid.

COMPOUND OF CARBON WITH NITROGEN.

Of Bicarburet of Nitrogen, or Cyanogen.

1295. Cyanogen ranks next to iodine among electrogative bodies. It is included among the halogen bodies Berzelius, and in the basacigen class by me. (625, 634.) ing a compound, I have deferred treating of it until w.

1296. Preparation.—Cyanogen is obtained by subjecting re and dry bicyanide of mercury to a low red-heat in a reclain or coated glass retort or tube, and receiving the

duct over mercury.

1297. Properties.—Cyanogen is a colourless, transpath, irrespirable gas, which painfully affects the nose and es, and has a strong and peculiar odour. Under a presse of four atmospheres, it becomes a colourless liquid, her than water. It may likewise be liquefied, or even idified by cold. It is characterized by burning with a autiful violet flame. It is decomposed by the electric ark, or by an incandescent iron into its constituents, carn and nitrogen. Alcohol dissolves twenty-three times, d water four and a half times its volume of cyanogen. In course of a few days the solutions become discoloured, d a brown matter is deposited. The deposition from al-

cohol has been found to contain carbon and nitrogen. After obtaining cyanogen from the bicyanide of mercury, a black residuum is found in the retort, which has been conceived to consist of carbon with a lesser proportion of nitrogen than exists in cyanogen; but of late, this residuum, and the deposition from alcohol, have been supposed to be isomeric with cyanogen.

1298. When ignited with two volumes of oxygen, a volume of cyanogen is converted into two volumes of carbonic acid and one of nitrogen, without condensation. Of course, as each volume of carbonic acid requires a volume of carbon vapour, there must exist two such volumes in one of cyanogen. Hence, as in the case of carbon and nitrogen each volume represents an atom, cyanogen consists of

two atoms of carbon and one of nitrogen

= 12= 14

and its equivalent is

26

Of the Nomenclature of the Compounds of Cyanogen.

1299. When Prussian blue is digested with a solution of potash, and the resulting solution is filtered while hot, yellow crystals are deposited by refrigeration, called ferroprussiate or ferrocyanate of potash, under the idea that they consist of an acid composed of iron, cyanogen, and hydrogen, in union with the oxide of potassium. Berzelius considers these yellow crystals as a double salt, formed by a "cyanure" of iron, and a "cyanure" of potassium. The name of this double salt, agreeably to his nomenclature is "cyanure ferroso-potassique." There is another compound containing the same elements, in which the proportion of cyanogen to that in the first mentioned compound, is as 1½ to 1, and for which his name is "cyanure ferrico-potassique."

1300. The existence of these combinations constitutes one instance among many, in which, according to Berzelius, two compounds, each having the same halogen body as an ingredient, form by their union a double salt.

1301. Agreeably to his system, we have double "chlorures, bromures.

fluorures," and "iodures," as well as double "cyanures."

1302. Some years ago, Bonsdorf, a skilful and sagacious German clemist, assailed this classification of Berzelius, by showing that some of the "chlorures" of the double salts exercised an alkaline, others an acid rection, with vegetable colouring matter; and that consequently the double "chlorures," so called by Berzelius, were really simple salts, in which on chlorure acted the part of an acid, the other of a base. Merely on contemplating the facts of the case, as stated by Berzelius, without have any knowledge of Bonsdorf's experiments and conclusions, the convictor arose in my mind that the double haloid salts, of that great chemist, show be considered as compounded of acids and bases. I cannot conceive where fore Bonsdorf thought it necessary to show that the ingredients of a double chlorure should be capable of reacting with vegetable colouring matter, as a

acid, the other a base, in order to prove their pretene of them were as severally, to acidity and basidity. (629.) It appears to me, that, cepting in the case of the alkalies and alkaline earths, those properties we not been deemed essential to oxacids and oxibases, and that of course ey ought not to be required in acids or bases formed by any other of the sacigen class. Agreeably to the definition of acids and bases, on which e basacigen classification is founded, (6: 10 632,) the "cyanure" of iron ing electro-negative as contrasted with "cyanure" of potassium, the e must be deemed a cyanobase, t r a cyanacid. 1303. It has been mentioned the DV British and French chemists the a compound formed by a supporter mination in ide was made to ind , while the termination in wret combustion with a combustible or pound formed of two radicals. wes was employed to designa 8. e difference in the practice of the two 100ls arose from the extension the class of supporters by the cner of Great Britain to the simple logen elements of Berzelius, who are according to those of France, oxygen as the only supporter, all the other elements being combustibles or radils. (685, &c.) Hence, according to the latter, only the compounds formed roxygen have been distinguished by the termination in ide as in oxide; hile, according to the former, in addition to those formed with oxygen, we re such as are formed by chlorine, bromine, iodine, and fluorine, distinished by the termination in ide, as has been already, to a certain extent, plained. (685.)

1304. By Berzelius the termination in ide is only resorted to where the dical is an electro-negative body; or, in other words, a body of which exides go to the positive pole. When the radical is one of those bosowted, when oxydized, go to the negative pole, the termination in we is sorted to. I object to this complicated nomenclature, as founded on the ror of not allowing those characteristics of acids and bases which have en acted upon by chemists in general, and by Berzelius himself in the se of oxacids and oxibases, to extend to the binary compounds formed by

e bodies of the halogen class.

1305. I consider the yellow salt in question, as consisting of a cyanacid maining an atom of cyanogen and an atom of iron, and which I would ll cyanoferrous acid, united to a cyanobase of potassium, consisting of e atom of cyanogen, and one atom of potassium, and forming a cyanorite of potassium. The double salt, consisting of the same elements, at containing both in the acid and base, half an atom more of cyanogen, buld, by analogy with the oxacids, have its acid distinguished by the time of cyanoferric acid, and should itself be called cyanoferrate of potassium.

Of Cyanic, Cyanuric, and Fulminic Acids.

1306. An atom of cyanogen, combined with an atom of oxygen, forms maic acid, which may be obtained in union with potash, by igniting raide of manganese with ferroprussiate of potash, or cyanoferrite of tanium; being the salt alluded to above, as consisting of cyanogen, iron, ad patasium. The cyanogen and potassium are converted, by the excess oxygen in the manganese, into cyanic acid and potash, which unite, raing a cyanate of potash. Cyanic acid cannot, however, be obtained one the cyanates, in consequence of its extreme susceptibility of decom-

1307. A crystalline substance may be procured from human urine, which

is known by the name of urea. It co sof bon, 1 rogen, oxyge and hydrogen, in the proportion to form one atom of cyanic acid, one atom of ammonia, and one atom of water. When urea is subjected to hat, ammonia escapes, and an acid remains, which was supposed to consist of one atom of cyanogen, and two atoms of oxygen. But it has been recently ascertained by Wohler and Liebig, that it consists of the elements of cyanis acid, chemically united to the elements or water; an atom of hydrogen, and an additional atom of oxygen, entering into its composition, not as water, but as essential constituents. Under these impressions, a new name, cyanuric, was given to it. This acid is solid, fixed, inodorous, and nearly tasteless. By combining with two atoms of water, as water of crystallization, it becomes capable of forming large crystals.

1808. When anhydrous cyanuric acid is exposed, in a glass retost, to a low red-heat, the extricated vapours being collected in a receiver refrigerated by a freezing mixture, hydrous cyanic acid is obtained. This acid and cyanuric acid consist of the same elements in the same proportion, but possess different properties and atomic weights. Hydrous cyanic acid is a colourless, volatile liquid, possessing a penetrating odour resembling that of acetic acid. It vesicates the skin when applied to it, exciting intense pain. Its vapour reddens litmus paper, is inflammable, and so pungent as to produce tears, and cause severe pain in the hands. Cyanuric acid is comparatively inert in these respects, but is far less susceptible of decomposition; as it is not decomposed by solution in boiling nitric or sulphants acid, while hydrous cyanic acid is decomposed by the addition of water-

1809. Hydrous cyanic acid, at the ordinary temperature of the air, spontaneously undergoes an explosive decomposition, attended by an evalution of heat, and is converted into a solid mass of dazzling whiteness. This mass consists of a variety of cyanuric acid, which differs from that shove described, in being insoluble in water or nitric acid, and in being decomposed by sulphuric acid. It is, therefore, to be considered as presenting a case of isomerism. (1153).

1310. It is remarkable that, although cyanuric acid consists of the same elements in the same proportion as hydrous cyanic acid, it carries the hydrogen and oxygen which exist in it in the proportion to form water, into every combination which it forms; while the hydrous cyanic acid, in combining with bases, separates from the water, which must be considered, when in union with this acid, as acting as a base.

1311. To bodies which, although they contain the same elements in the same ratio, yet hold them differently associated, so that in reacting with other agents, they are resolved into, or form compounds differing in composition, the term metameric has been applied. Thus hydrous cyanic, and

cyanuric acid are said to be metameric with regard to each other.

1312. Another compound of cyanogen with oxygen exists in the fulminating mercury of Howard, and the analogous fulminating silver of Desco-Liebig ascertained these compounds to contain an acid common to both, which he called fulminic acid, but which, agreeably to the analysis made by him and Gay Lussac, was identified in composition with cyanic acid. Yet, as the latter would not produce fulminating compounds, and differed in its other properties, these acids have been considered as affording another instance of isomerism. Mr. Edmund Davy, however, alleges the existence of hydrogen in fulminic acid, and likewise that the nitrogen exists in excess, beyond the proportion appropriate to evanogen.

1313. Fulminic acid is a colourless, transparent, volatile liquid, which

carbon. 245

s litmus, and produces a taste at first sweet, but afterwards astringent sagreeable. Its fumes have a pungent and disagreeable odour, and he headach when incautiously inhaled.

4. Besides these acids, M. Liebig has recently discovered another, is polymeric with regard to cyanuric acid; as it consists of the same ts in the same ratio, though twice as much of each enters into the sition of an atom.

Of the Chlorides, Bromides, and Iodides of Cyanogen.

5. Chlorine forms two compounds with cyanogen, a protochloride perchloride. The *protochloride* is a colourless, fetid gas, which may refied, and even solidified by cold. In common with several other ands of cyanogen, it possesses, even when gaseous, the singular proof producing pain by contact with the skin. The *perchloride* is a crystalline substance, with an odour resembling that of mice.

6. Bromine and iodine both form with cyanogen, crystalline com-

3. The chlorides and bromides of cyanogen are energetic poisons.

Of Sulphocyanogen.

7. It has been stated that the yellow salt, usually known as ferrote of potash, is by Berzelius considered, when free from water, as ing of cyanogen, iron, and potassium; also that I consider it as a errite of the cyanobase of potassium. When this salt, desiccated to cence and finely pulverized, is mingled with flowers of sulphur, and d to a red-heat in a porcelain crucible, the iron is displaced; the sulnid cyanogen uniting, form a compound called sulphocyanogen, and niting with the potassium, constitutes a sulphocyanide. (1302.)

8. Sulphocyanogen has been isolated by passing chlorine through a in of sulphocyanide of potassium, or by subjecting that compound to acid. Sulphocyanogen has some pretensions to be classed with the

n, and of course with the basacigen bodies.

9. The intense blood-red colour which it produces with iron, is the triking property of sulphocyanogen, and has led to the impression e sulphocyanide of iron may be the colouring matter of the blood.

0. Sulphocyanogen is solid, insoluble in water or alcohol, and may, inhydrous state, be sublimed without change. It is composed of one

of cyanogen, and two atoms of sulphur.

1. Dr. Thomson states that another compound of sulphur and cyanotists, containing one atom of sulphur and two atoms of cyanogen. ompound may be obtained in transparent colourless crystals. It is possesses a strong smell, and is soluble in water. When applied tongue, even in a minute quantity, it produces intense pain; and the uched remains red and painful for some time.

Of Sulphocyanhydric Acid.

2. This acid may be obtained from a solution of the sulphocyanide tassium, by the addition of phosphoric acid. Water is decomposed, tygen unites with the potassium, forming potash, with which the phosphoric acid combines, and the hydrogen with the sulphocyanogen, forming tryanhydric acid, which may be separated by distillation. This acid and colourless, has an acid taste, and powerful odour. It becomes

solid at 14°, and boils at 216°. It is composed of one storn of sulphe-cyanogen, and one atom of hydrogen.

Of Cyanhydric or Prussic Acid.

1323. One atom of cyanogen, equivalent 26, with one atom of hydrogen, equivalent 1, forms one atom of cyan-

hydric acid, equivalent 27.

1324. This acid has been detected in water distilled from bitter almonds and from laurel leaves, also, from peach leaves or blossoms. Between the odour of these and that of the acid when dilute, it would be difficult to discriminate.

1325. Laurel water has long been known as a poison Water distilled from peach leaves has been used to impart an agreeable flavour to food. Some peach leaf water prepared by Mr. Wetherill, gave indications of cyanhydriacid, by producing a blue colour with a solution of iron.

1326. There have been instances in which noyeau, a cordial made from the kernels of bitter almonds, had proved poisonous from the presence of cyanhydric acid.

1327. There is a salt consisting of two atoms of cyanogen and one of mercury, called bicyanide of mercury. When this salt is subjected to the action of chlorohydric acid, the chlorine forms a chloride with the mercury, while the hydrogen forms cyanhydric acid with the cyanogen.

1328. It may be more conveniently obtained by impregnating with sulphydric acid, a solution containing sixty grains of bicyanide of mercury for every ounce of water. The hydrogen unites with the cyanogen, while the sulphur precipitates with the metal. Any excess of the sulphydric acid is easily removed by the carbonate of lead. The apparatus for impregnation with sulphydric acid, has been described already. (797-8.)

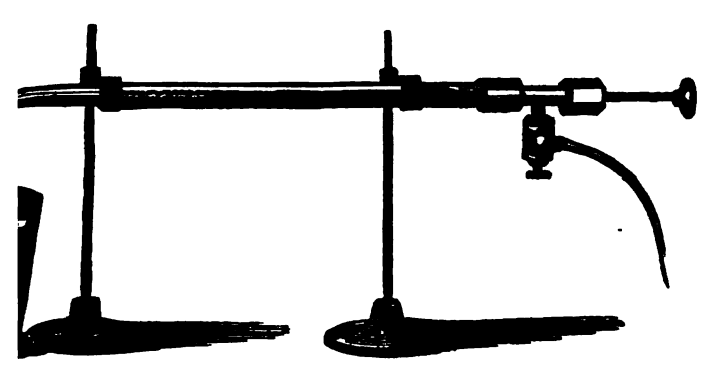
1329. The acid may be procured in its most concentrated form, by exposing the bicyanide in crystals, in tube, to sulphydric acid gas, and employing a receiver, surrounded by salt and snow, to condense the vapour

evolved.

1330. In performing this process, I found great difficulty to arise from the inability of the operator to regulate the quantity of gas introduced into the tube, so that, on the one hand, there might be no absorption of atmospheric air, and, on the other, no excess of the gas escaping, and

puently causing a loss of materials, and annoyance bystanders. This difficulty is in great measure rel, by means of the apparatus of which an engraving escription is subjoined.

Apparatus for the Evolution of Cyanhydric or Prussic Acid.



set a tube, three-fourths of an inch in bore and about two feet in length, be ight angles, at about six inches distance from one end. Let the shorter drawn out into a tapering form, with a bore not exceeding a tenth of an ameter. Upon the larger orifice let a brass band be cemented, in which a rew has been cut, so that a stuffing-box, furnished with a corresponding w, may be easily fastened air-tight to the band, or removed when desirable. the stuffing-box an iron rod passes, flattened like an oar at the end, which the tube when the stuffing-box is in its place. There must likewise be a perture in the band communicating with the cavity of the tube, and furith a gallows screw. The main body of the tube is to be situated nearly a little inclined towards the curvature, so that the tapering extremity may mearly perpendicularly into a tall narrow phial, surrounded by a freezing The horizontal portion of the tube near the bend should likewise be re-The apparatus being thus arranged, introduce a sufficient quantity of inide of mercury into the tube, and close it by inserting the stuffing-box ed. In the next place, by means of the gallows screw, make a communiween the cavity of the tube, and a self-regulating reservoir of sulphydric is gas must be allowed to pass into the tube very slowly, and meanwhile, of the rod, the bicyanide is to be stirred. Before long a portion of the cyacid will be seen in the narrow part of the tube. This serves to regulate ssion of the sulphydric acid, since, when the quantity passing into the tube rate, the liquid will rise in the tube; when too great, it will be expelled By these means, after a little while, all the bicyanide will be decomposed, responding quantity of acid collected in the refrigerated phial. lince this figure was engraved, I have found it preferable to have a phial 1 a bottom tapering to a point, so that the quantity of acid, however minute, ipperent; and it is sooner rendered competent to act as an index of the prohe process; so as to regulate the quantity of gas to be allowed to enter the has also been found advantageous to mix the bicyanide intimately with e its bulk of glass, powdered to the consistency of coarse sand.

New Process for Liquid Cyanhydric Acid.

The following process for procuring prussic acid, is recommended toor Everitt.*

For every 212 grains of ferroprussiate of potash (cyanoferrite of m,) in 2 ounces of water introduced into a retort, add as much sul-

• Lendon and Edinburg Philosophical Magazine, vol. 6, p. 100.

phuric acid as may be equivalent to 120 grains of the anh drous acid; and distilling the mixture, let the vapour pass into a pint of resigning rated water, holding 255 grains of nitrate of silver. The resulting precipitate being washed and dried, should constitute nearly 201 grains of mercurial cyanide. Of this let 40 grains be introduced into 7 fluid ounces, and 20 minims of water; and add 40 minims of chlorohydric acid, of specific gravity of 1.12. The whole being well secured in a stoppered bottle, and agitated repeatedly, should be allowed to rest until the resulting chloride of silver subsides. In the solution thus obtained, when carefully decanted, there will be one grain of prussic acid (more properly called cyanhydric acid,) for every fait ounce of water.

1335. Should there be a little excess of chlorohydric acid, agreeably to Professor Everitt's observation, confirmed by those of others, it will tend rather to preserve, than to decompose the acid.

1336. Properties of Cyanhydric Acid.—This acid is a colourless liquid, which emits a powerful odour, resembling that of peach blossoms. When perfectly free from water, it is far more volatile than ether, as it boils at 79° F., and evaporates so rapidly, that one portion becomes frozen by the loss of the caloric which the other absorbs in passing into the aëriform state. Its specific gravity is 0.7058, be-

ing nearly the same as that of sulphuric ether.

1337. Anhydrous cyanhydric acid is sometimes decomposed in a few hours, especially if not protected from the light, and can never be preserved longer than a fortnight. Either when in the state of a liquid, or vapour, this acid is probably the most active poison known. The application of a few drops to the arm of a man has produced death, and its fumes are equally deleterious when inspired. As when free from water, this acid boils at 79°, nearly 20° below the temperature of the blood, it must be converted into vapour too soon to produce its full effect. From a cavity like the ear, the pure acid must be ejected in vapour immediately. I am, therefore, under the impression that it is less effectual as a poison when anhydrous, than when combined with a minute proportion of water.

1338. Upon one occasion, touching the ear of a rat confined in a glass jar with a drop of the anhydrous acid, the animal, being obliged to breathe the vapour, died instantaneously with a slight sneezing. Yet upon another occasion nearly half a drachm was injected into the ear of a large dog, without causing death; a like quantity, subsequently injected into his nose, proved fatal. The acid earployed was so pure as to freeze by its own evaporation.

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1339. The best antidotes for this poison are chlorine or amonia, in dilute aqueous solution, especially chlorine.

- 1340. Cyanhydric acid is sometimes employed in medine, though in very small doses, and in a very diluted ite.
- 1341. It has been proposed to detect cyanhydric acid, cases in which it may have been employed in poisoning, subjecting the stomach and its contents to distillation th water, and testing the liquid product by copper or n.
- 1342. I should place much reliance on the characteristic ell of the acid, which is that of peach blossoms, and ich may be perceived, not only from the presence of e acid, but likewise from that of any of the cyanides, if bjected to the action of chlorohydric acid.

Experimental Illustrations.

1343. The processes for the production both of the neous and anhydrous cyanhydric acid, exhibited; also, congelation of the latter by the cold arising from its on evaporation.

SECTION V.

OF BORON.

1344. Preparation.—By the addition of sulphuric acid a saturated solution of biborate of soda (borax) in war, shining crystalline plates are precipitated, consisting of ric acid. From these crystals boron may be obtained, ther by the action of a powerful Voltaic series, or by ret vitrifying them, then finely pulverizing the resulting lass, and afterwards heating the acid thus prepared in patact with potassium.

1345. Boron may be obtained by means of the appartus employed for the evolution of silicon, (1355, &c. 357, &c.) substituting fluoboric acid gas for fluosilicic

cid gas.

1346. Properties.—Boron is of a dark olive colour, tastess, inodorous, a non-conductor of electricity, and insolue either in alcohol, ether, or the oils. Its atomic weight

When heated in the air to 600° F, it takes fire, and, by uniting with oxygen, generates boric acid. Nevertheless only a portion of the boron is oxydized, the remainder being protected by a crust of fused boric acid. If this crust be removed by water, the boron will be found to have undergone a change similar to that produced in charcoal by an intensely high temperature. It is rendered harder, more difficult to ignite, and so much denser, that, although its specific gravity was before only 1.83, it now sinks rapidly in sulphuric acid of the specific gravity of 1.844. Before it has been ignited, boron is slightly soluble in water; and its solution, when evaporated to a certain point, forms a gelatinous mass, which, by complete desiccation, becomes opaque, and assumes the usual appearance of boron.

COMPOUND OF BORON WITH OXYGEN.

Of Boric* or Boracic Acid.

1347. The means of procuring this acid have been mentioned in describing the process for obtaining boron. Borax is a biborate of soda, from which boric acid may be liberated in crystals, as above described, by the superior

affinity of sulphuric acid for the soda.

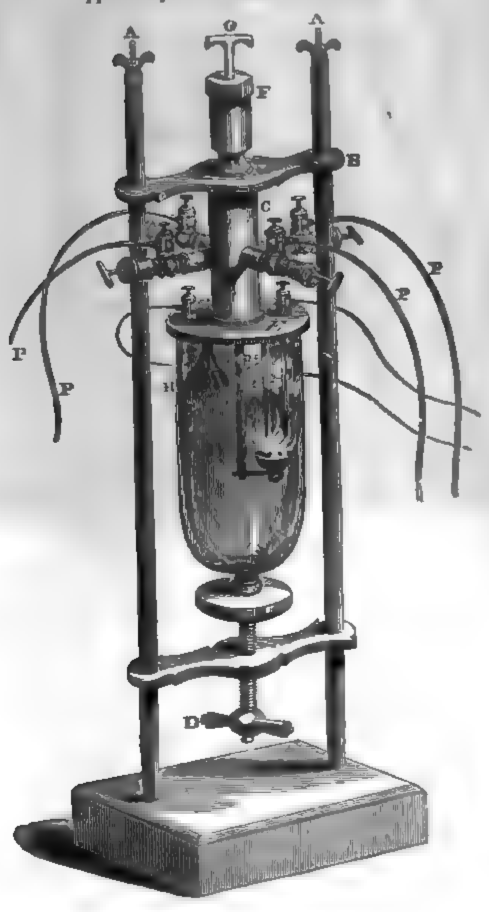
1348. Properties.—Boric acid is crystalline as first obtained from borax, but forms a glass when deprived by heat of its water of crystallization. It is colourless, inodorous, almost tasteless, and sparingly soluble in water. In the form of an aqueous solution, its agency is weak, and it is in consequence rarely used in that state. In common with silicic, phosphoric, and arsenic acid, being fixed at temperatures at which sulphuric and nitric acid are decomposed, it will at those heats expel them from their combinations; although, when water is present, and at low temperatures, it is displaced from combination not only by those acids, but by many others. It consists of one atom of boron, and three of oxygen.

1349. Boron, in its habitudes, seems to lie between phosphorus and carbon. In its insusceptibility of volatili-

^{*} I agree with the French chemists and Berzelius, in employing the word levic instead of boracic, as more naturally generated from boron, by analogy with the other acids formed with radicals, to the last letter of which the letters ic are usually added.



Apparatus for the Evolution of Silicon.



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tion, infusibility, and the temperature requisite for its mbustion, it is most allied to carbon; yet boric acid is ore analogous to phosphoric than to carbonic acid. oth phosphoric and boric acid are capable of being reced to a vitreous state, and bear a white-heat without ing volatilized; while the acid of carbon is naturally riform.

1350. Boric acid and the biborate of soda are of great in blowpipe assays, as fluxes, and in soldering, as the ans of protecting metallic surfaces from oxidation.

Experimental Illustrations.

1351. Saturated solution of borax, decomposed by suluric acid. Exhibition of crystals of the acid and of the orate, which are severally fused into a glass by the npound blowpipe. Effects of cobalt and manganese on the colour of the glass, of which a globule is convently supported by a platinum wire.

Of Chloride of Boron.

1352. The chloride of boron may be obtained by the combustion of boron chlorine; or by passing a current of chlorine over a mixture of charcoal d boric acid, heated to redness in a porcelain tube.

1353. The chloride of boron is a colourless gas, possessing a strong and suliar smell. When brought in contact with water, a reciprocal decomition takes place, and boric and chlorohydric acid result. It forms a ite salt with ammonia, and is by some chemists considered as an acid.

SECTION VI.

-m+0 **(1)** 0+m--

OF SILICON.

1354. Preparation.—By heating sulphuric acid with a sture of powdered Derbyshire spar, and powdered glass, quartz, a permanent gas may be obtained. When possium is heated in this gas, silicon is evolved.

Apparatus for evolving Silicon from Fluosilicic Acid Gas by means of Potassium.

This apparatus is represented by the opposite engraving. Into a stout marray block as a basis, two iron rods, A A, are so planted as to extend perpendiculty, and of course parallel to each other, about two feet in height. Upon these is two iron bars are supported horizontally, one B, near their upper extremities, other, at the height of about six inches from the wooden basis. In the centre of lower bar, there is a screw, D, having a handle below the bar, and supporting we it a circular wooden block. Into a hole in the upper iron bar, equidistant from reds, is inserted a hollow brass cylinder, C, which at the lower end screws into

an aperture in a circular plate of brases, E, which is thus supported horizonly fow inches below the bar. By these means, room is allowed for the insertion the cylinder of four valve rocks, each furnished with a gallows screw. The cyl is surmounted by a stuffing hox. F. through which a copper sliding rod, G. and aght. The breez plate is turned and ground to fit a bell glass of about five it in diameter, and eight meters in height, which is pressed up when necessary but the plate and the 11 ack, by the serew, D, supporting the block. Within the comprised by the bell glass, and on one side of the centre of the plate, two brass wires are inscribed, one of them insulated by a collar of leathers, so as to of the ignition, by a galvanic discharge, of a small arch of platinum wire, a reaches from one to the other. The sliding-red abovementioned as occupying stuffing box terminates below the plate in an elbow which supports a cup at angles to the rod, at the same distance from the rod as the platinum wire, as the opposite aids at it, there is a brain cover, II, for the cup, supported from plate. The arrangement is such, that by a suitable movement in the sliding made by grasping it by the handle, G, in which it terminates externally, the may be made either to receive into its cavity the platinum wire, or to adjust its

its cover, H
1356. The bell being removed, about sixty grains of polasions, in pieces ast
taining more than fifteen grains each, are to be introduced into the cup, whi then to be adjusted to the cover, and the bell secured. In the next place, by a of the flexible lead tubes, P. P. P. P. and the gallows screws attached to the cocks, established a communication severally with an air pump, a self regal reservoir of hydrogen, a barometer gauge, and a jur over the mercurial custom, tuning fluosilicie acid gas. First, by means of the air pump, exhaust the bell in order to wash out all remains of atmospheric air, admit by dregen from the twoir. Again exhaust, and again admit hydrogen. Lastly, exhaust the bell of higher, and admit the fluosilicie acid gas. By means of the gauge, the exhaust indicated and measured, and by the same means it will be seen when the promittee are within the bell approaches that of the atmosphere. When this takes the gas within the bell approaches that of the atmosphere. When this taken? the cocks being all closed, and by means of the process of galvano-ignition, &c.) the platinum wire being rendered incandescent, the polassium is to be be into contact with it. A poculiar deep red combustion ensues, evolving out chocolate-coloured fumos, which condensing into flocks of the same hue, de throughout the receiver, and are deposited upon the interior surface, so as to \$ in the mind of the spectator, the idea of a miniature full of chocolate-coloured i On removing the bell after the potassium has ceased to burn, the cup which h in found to contain silicon mixed with the fluoride of potassium, and with the whole of the chocolate-coloured deposition is contaminated. Silicuiret of potas to bkewise found in the cup, since, upon the affinion of water, a fetid inflam. gas is evolved, which has an edear resembling that of pleaspheretted hydrogen. which must obviously be the analogous compound afficiaretted hydrogen con, being insoluble, may be separated from the fluoride by digestion in water V the potassium employed is of the kind obtained by means of charcoal, the silice as Berzelius alleges, adulterated with carbon. I am under the impression that si nitric acid removes this impurity.

Simple Process for the Evolution of Silicon.

1357. Last winter I was enabled to adopt a much more simple and: venient process for the evolution of silicon, which is as follows:

1358. A bell glass was filled, over mercury, with fluosilicic acidmeans of a bent wire a cylindrical cage of wire-gauze, containing a globules of potassium, was introduced through the mercury into the co of the bell, and supported in a central position. A knob of iron was we to the end of a rod, of the same metal, so recurved as to reach the with case. Having been heated nearly white-hot, this knob was pe through the mercury, so as to touch the cage. By these means the p sium having been made to enter into combustion with the fluorine, the con was evolved. Much of this substance remained attached to the cap combination with fluoride of potassium. From the impurities, with w it was thus associated, the silicon was separated by washing in water digestion with nitric acid. There can be no doubt that this process

he employed to evolve boron, by employing fluoboric acid instead of fluo-silicic acid.

- 1359. Properties of Silicon.—It is of a brown-ash colour, without the least trace of metallic lustre, a nonconductor of electricity, infusible, and incapable of being rolatilized. It is not liable to be dissolved or oxydized by rulphuric, nitric, chlorohydric, or fluohydric acid, but is roluble in a mixture of nitric and fluohydric acid. When beated to redness with the fixed alkaline carbonates, it burns vividly; and when dropped upon the hydrates of potash, soda, or baryta, while in a state of fusion, it explodes. Yet it is unchanged by ignition with chlorate of potash, and exercises but a feeble reaction with nitre, even when heated to redness. In these respects its habitudes are anomalous.
- 1360. When silicon, as usually obtained by the aid of potassium, is intensely heated in the air or in oxygen gas, it burns with a feeble blue flame; but, by becoming encrusted with silicic acid, a portion escapes combustion. This portion is rendered harder, denser, and insusceptible of combustion with oxygen at the highest temperatures. Berzelius suspects the greater combustibility, and inferior density and hardness of silicon, in the state in which it is obtained by the process above described, to be due to the presence of hydrogen, derived from the water employed. In this state, it inflames when ignited in the vapour of sulphur, and forms a sulphide, which is decomposed by water into sulphydric and silicic acid.

COMPOUND OF SILICON WITH OXYGEN.

Of Silica, or Silicic Acid.

- 1361. One atom of silicon with one atom of oxygen, each equivalent to 8, forms one atom of silicic acid, equivalent 16.
- 1362. Preparation.—Quartz being powdered, and fused with three times its weight of pearlash, a glass is obtained, which, being soluble, forms with water a liquid, called formerly liquor silicum, or liquor of flints. An acid being poured into this solution, silicic acid, slightly contaminated by potash, is precipitated.

1363. To obtain silicic acid, Berzelius advises us to fuse in a platinum crucible, equal parts of the carbonates of

potash and soda, and to add quartz, finely ulverized, in small successive portions. The effervescence arising from the addition of one portion, is allowed to subside before adding another, until effervescence can no longer be excited. The refrigerated mass is dissolved in chlorohydric acid, and the solution filtered and evaporated to dryness. To remove all traces of iron or alumina, the dry mass is kept moist with chlorohydric acid, during about two hours, and afterwards washed with hot water, and then exposed to a red-heat. Silicic acid will remain in a sufficient degree of purity.

1364. Pure silicic acid, in the well known form of rock crystal, is found throughout nature. Its usual crystaline form is a six-sided prism, terminated by a pyramid with

six faces.

1365. Properties.—Pure silicic acid is white, tasteless, and inodorous, and has a specific gravity of 2.66. Its solution does not redden litmus, and, when evaporated to a certain point, forms a translucent jelly. It is soluble when nascent, but insoluble after exposure to heat or desiccation, or in its native crystalline form.

1366. It was first fused by myself, in the year 1801, by means of the compound blowpipe. It has never been

volatilized.

Of Chloride of Silicon.

1367. When silicon is heated in chlorine it inflames, evolving heat and light, and a chloride of silicon is formed, which is a volatile liquid, possessing a sharp and powerful odour. In consequence of the absorption of an excess of chlorine, it is generally coloured yellow. It boils below 212°, and, by the addition of water, is converted into chlorohydric and silicic acid.

Experimental Illustrations.

1368. Silicate of potash, exhibited; also the solution of it, called liquor silicum, from which silica is precipitated by means of an acid.

Of Glass.

1369. If the proportions, in which sand and alkali are used as above mentioned for the liquor silicum, be reversed, the insoluble compound of silicic acid and alkali, known under the name of glass, is obtained, which however pure the materials, has a slight tinge of green. This is removed by a due admixture of the red oxide of lead, and black oxide of mangunes 1370. Annealing Process.—A sudden diminution of the quantity of

silicon. 255

c among the exterior particles of a thick piece of glass in a state of on, is not attended by a corresponding diminution of the quantity of rinciple among the particles within, owing to the slowness with which conducts heat. Hence, there can neither be a general coherence, nor form arrangement among the particles; unless the cooling be very so as to allow the refrigeration, within and without, to be nearly sineous. As it never can be perfectly simultaneous, the annealing ways be defective, other things being equal, in proportion as the glass ker. Were the particles subjected to radiant heat only, the process be more effectual; as this, when proceeding from incandescent surhas been ascertained to penetrate and even to pass through glass.

11. By gradually making up a fire of charcoal, at about four inches ce on each side of a glass tube of about an inch and a quarter in ess, and with a very small bore, I was enabled to heat it red-hot, at causing a fracture. From its situation, it was subjected to radiant

mly.

12. By opening a perpendicular hole in an anthracite fire, I have been id, with little delay, to introduce the beaks of glass retorts of two or gallon in capacity, without causing a fracture. Thus situated, the soon becomes almost fluid, so that by its own weight the lower pordrawn downwards into a tapering tube, and would be made to fall the the beak not removed from the fire. If removed in due time, the of the retort may be so held as to cause the tapering portion of the oform such an angle with the other part, as to be capable of entere tubulure of another retort, as described in one of the processes for ring pure chlorohydric acid. (891.)

73. By like means, the beaks of broken retorts, or any piece of a glass may be made to taper, to be elongated so as to be inserted through bulure of a retort, and to serve, consequently, when luted to the tubufor the introduction of sulphuric acid, in various processes besides that

ich allusion is above made.

14. Prince Rupert's Drops.—When glass, in a state of fusion, is ed into water, the defective states of cohesion and arrangement, connat to the want of annealing, are at the maximum. Such drops have seen known under the name of Prince Rupert's drops. It is only sary to break off the slender filament in which the mass terminates, ler to cause an explosive dispersion of the whole into a coarse powder. 75. The cohesion of the particles in glass tubes, is often nearly as instant in Prince Rupert's drops. The slightest mark from a file on the present of the prince representation of the prince surface, or even wiping them out, especially if a metallic wire be sped, may cause them to break into pieces. Sometimes the fracture immediately, at other times, not till many hours have intervened.

MPOUNDS OF FLUORINE WITH HYDROGEN, BORON, AND SILICON.

176. Fluorine has been briefly noticed, (746, &c.) I med treating of the interesting compounds formed by element with hydrogen, boron, and silicon, until the ent should be acquainted with those substances.

177. The three fluorides referred to are called sevefluohydric, fluoboric, and fluosilicic acid. (862).

Of Fluohydric Acid, generally called Hydrofluoric Acid. (850.)

1878. Fluorine exists in nature in union with the metals of the carts and alkalies, especially with calcium, a metal of which lime is the oxide. Such compounds are called fluorides. The remarkable mineral, called

Derbyshire or fluor spar, is a fluoride of calcium.

1879. Not long since, Derbyshire spar was considered a compound of lime with an acid, called fluoric acid, and supposed to consist of oxygen and an unknown radical. Mr. Ampere first suggested the present doctrine, which was soon adopted by Sir H. Davy, and is now, I believe, universally sanctioned.

1380. Preparation.—When fluoride of calcium is pulverized, and hated in a leaden retort with twice its weight of concentrated sulphuric acid, the water in combination with the acid is decomposed. The oxygen and acid form sulphate of lime with the calcium; while the hydrogen produces with the fluorine, fluohydric acid, which passes over in the form of a very wintile acid vapour, and may be condensed in a leaden or silver receiver, servounded by a mixture of snow and salt. If received in water, it condenses

without refrigeration, and forms a diluted acid.

1881. Properties.—Fluohydric acid is a colourless, limpid liquid, which boils at a little below 60°. When anhydrous, its specific gravity is 1.0666.

It is so volatile, that, in a close apartment it cannot be decanted without subjecting the operator to intolerable fumes. This operation must be

formed where there is a current of air to carry them off.

1382. It ulcerates the skin with peculiar activity, and corrodes glass so as to trace its course indelibly, in running over the surface. It must be kept in vessels of silver or lead, accurately closed. When received in water it is absorbed, forming aqueous fluohydric acid, and is then more easily preserved.

1383. One atom of hydrogen, equivalent 1, with one atom of fluorise, equivalent 18, is supposed to form one atom of fluohydric acid, equiva-

lent 19.

Experimental Illustrations.

1384. Powdered fluoride of calcium, heated with suphuric acid in a leaden retort, adapted to a receiver surrounded by a mixture of snow and salt. Same process, substituting a receiver with water, by means of Knight's apparatus. Effect of fluohydric acid upon glass.

Of Fluoboric Acid.

of two parts of powdered fluoride of calcium, with one of vitrified bore acid, in an iron tube. One part of the boric acid is decomposed, the oxygen of which, and the remaining portion of the acid, form borate of lime with the calcium; while the boron unites with the fluorine, forming fluoric acid gas, which must be received over mercury. Fluoboric acid gas, may likewise be procured, by heating in a glass retort two parts of fluoride of calcium and one of boric acid, with twelve parts of concentrated sulpharic acid. Berzelius, however, states that, when obtained by this method, it is contaminated by fluosilic acid, arising from the action of the fluorine on

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This might, however, be avoided by performing the operation den retort.

7. Dr. Thomson states that the best method of obtaining fluoboric is, is one which was suggested by Berzelius. Boric acid is to be in anhydrous fluohydric acid, and a gentle heat applied to the

A reciprocal decomposition takes place; the hydrogen of the ric acid combines with the oxygen of the boric acid, forming water, se fluorine unites with the boron, and constitutes fluoboric acid gas.

. Properties.—Fluoboric acid is a colourless, transparent gas, with destruc-Its specific gravity is 2.3622. Water absorbs seven hundred s volume of this gas. When fluoboric acid is passed into water, gen of a portion of the water unites with the boron, forming boric hile the hydrogen combines with the fluorine, producing fluohydric The boric acid precipitates, and the fluohydric acid combines with ecomposed portion of the fluoboric acid, forming a compound which is designates as hydrofluoboric acid, but which, according to the lature which I have adopted, should be called fluokydroboric acid. entinue to pass fluoboric acid gas into the water, or partially abstract id by evaporation, until the solution of fluoboric acid becomes satue affinities which were at first brought into play are reversed. The n of the fluohydric acid unites with the oxygen of the precipitated cid, and the fluorine with the boron; so that we finally obtain a solution of fluoboric acid in water. This solution is at first furning; the application of heat it yields up a fifth part of its gas, and then y resembles concentrated sulphuric acid in appearance. Like that carbonizes organic products, in consequence of its affinity for water. I. Three atoms of fluorine, equivalent 54, and one atom of boron, ent 11, form one atom of fluoboric acid, equivalent 65. (856, &c.)

Of Fluosilicic Acid.

Preparation.—It may be obtained by adding to the materials for g fluohydric acid, one half their weight of finely powdered glass, ng the mixture to heat in a glass retort, and receiving the product ercury; as by water it would be rapidly absorbed.

. The oxygen of the silicic acid in the glass, with the sulphuric acid cium, forms a sulphate of lime; while the fluorine and silicon escape

orm of fluosilicic acid gas.

Le The apparatus which I employ for fluosilicic acid, is precisely the sthat described under the head of ammonia.

Reporties.—Fluosilicic acid is a transparent, colourless gas, with iar and sufficating odour, closely resembling that of chlorohydric it reddens litmus paper, and has a specific gravity of 3.5735. When in contact with water it is rapidly absorbed, and a decomposition ace, similar to that which ensues in the case of fluoboric acid under circumstances. Silicic acid is deposited in the form of a gelatinous ad fluohydric acid is produced, which combines with the undecompation of the fluosilicic acid, forming a compound called hydrofluotid, to which, if it be an acid, I would give the name of fluohydrotic. If the water in combination with the fluohydrosilicic acid be ly removed by heat, fluosilicic acid gas escapes, leaving fluohydric solution.

1394. One atom of fluorine, equivalent 18, with one atom of sizes, equivalent 8, forms one atom of fluorilicic acid, equivalent 26.

Experimental Illustrations.

1395. Production of fluosilicic acid, shown: also its absorption by water, and the precipitation of silicic acid, as above described.

Of the Reaction of Fluohydric Acid with Fluoboric and Fluohicis Acid, and of the Nomenclature of the Compounds formed by the latter on meeting with Oxibases.

1396. The union which ensues between fluohydric acid, and either moboric, or fluosilicic acid, agreeably to the preceding statement, may appear anomalous, in the way in which it has hitherto been treated. If, however, I am correct in my mode of defining the difference between an acid and a base, (631,) the combinations in question will not prove to be anomalous. I deem it consistent to suppose that a fluobase of hydrogen unites, in the one case, with fluoboric acid, in the other with fluosilicic acid; so that hydroboric acid might be called fluoborate of the fluobase of hydrogen, acid would be called fluosilicate of the fluobase of hydrogen, or briefly.

fluosilicate of hydrogen.

1397. When either fluohydroboric acid, or fluohydrosilicic acid, er in other words the fluoborate or fluosilicate of the fluobase of hydrogen, is brought into contact with an oxibase, the radical of the latter takes the place of the hydrogen, which, with its oxygen, forms water. Thus, in the case of potash, there would result a fluobase of potassium, usurping the place of the fluobase of hydrogen; and of course either a fluosilicate, or fluoborate of potassium must be formed. Agreeably to the Berzelian nomenclature, these compounds are double salts, the name of one being in the French translation, "fluorure borico-potassique," that of the other, "fluorure silico-potassique." Many analogous salts, formed by the acids under consideration, with salifiable substances, are mentioned by Berzelius; also many others, in which other radicals, in union with fluorine, play a put analogous to that performed by silicon and boron, in the salts above meationed.

as acids. Of course it is consistent that compounds, usually called acids, should in some instances act as bases. In this respect, a striking analogy may be observed between the union of the oxide of hydrogen (water) with the oxacids and oxybases; and that of fluoride of hydrogen with fluoride and fluobases. According to Berzelius, water acts as a base to oxacide; as an acid to oxibases. So I conceive the fluoride of hydrogen acts as a base in the cases above noticed, while it acts as an acid in the compound of hydrogen, fluorine, and potassium, called by Berzelius "fluorure potassium acide." This compound I would call a fluohydrate of the fluobase of potassium, or more briefly, fluohydrate of potassium; as we say sulphate of copper, instead of the sulphate of the oxide (or oxybase) of copper.

SECTION VII.

OF ZIRCONION, OR ZIRCONIUM.

There is a stone, known under the name of the jargon or zircon on, from which Klaproth extricated an earth, to which the name of was given. This earth is an oxide of an elementary body, which n called zirconion, or zirconium. The termination in um being now plied by chemists to the names of substances having the metallic er, I think that it has been erroneously associated with the name of cent in question, since its pretensions to that character are not higher ose of carbon.

OF METALLIC RADICALS.

10. It is to metallic radicals that I deem it expedient next place to direct attention. Less than thirty ago, the line of demarcation between metals and bodies was easily drawn. There was then no known which had a specific gravity less than six; and of bodies, none of which the specific gravity was as as five. But the discovery of alkalifiable metallic als, having a specific gravity less than that of water, ilated the barrier which had been established on the

of superior gravity.

11. Peculiar brilliancy and opacity were in the next appealed to as the means of discrimination; and likethat superiority in the power of conducting heat and icity, which was so remarkable in substances of a edly metallic character. Yet so difficult has it been w the line between metallic and non-metallic radithat bodies which are by some authors placed in one are by others included in the other. Thus selenisilicon, and zirconion have by some chemists been rised among the metals, by others among non-metallic 3. In fact nature has not qualified her bodies for ct classification. It is true that there are those of the prominent features or qualities are so strikingly ent, that we are at first encouraged to think that by iating similar substances with each, we shall form es not liable to be confounded. Thus gold possesses, nigh degree, all the attributes of a metal, while sulphur ally devoid of them; yet arsenic, as being decidedly lic, may on the one side be classified with gold in

preference to sulphur; while on the other hand, between arsenic and sulphur, there is in many respects a much greater analogy than between arsenic and gold. In fact, tellurium, which had been always classified and is still considered as a metal, is now associated by Berzelius, in his amphigen class, with oxygen, selenium, and sulphur, and has, in consequence, been treated of by me as a basa-

cigen body.

1402. Metals were formerly distinguished as metals, and semi-metals; the latter appellation having been employed to designate such as were wanting in the mechanical properties of malleability and ductility. Again, the metals which were endowed with the properties just mentioned, were divided into noble and base. The noble metals, sometimes called precious, from their superior value, were distinguished from the others by their insusceptibility of injury from fire, moisture, or air. Silver and gold were, about a century ago, the only known metals meriting the name of noble, upon the grounds which I have mentioned To these platinum was subsequently added, and latterly palladium, nickel, iridium, and rhodium, have been found to have analogous pretensions, agreeably to the ideas in obedience to which the epithet was originally employed. Subsequently chemical properties became better known, and metals were associated not only in accordance with their own obvious characteristics, but also with a view to their oxides, which in many cases are the only forms under which they are met with in nature, or employed in the arts. Accordingly the metals are now generally divided with a view to their susceptibility of oxidizement, or the character of their oxides. Among the oxides alluded to there are some of which the characteristics are so different, that there can be no hesitation in classifying them separately. Yet in other members of the same class, the characteristics by which they are distinguished are so feeble, that a diversity of opinion has existed as to the genera to which they belong.

1403. I propose to divide metallic radicals into the for

following classes:

First, metals of the earths proper.

Second, metal's of the alkaline carths.

Third, metals of the alkalies, or alkalifiable metals.

Fourth, metals proper.

I shall employ the words noble to distinguish not liable to be tarnished by exposure to fire, or air; as, for instance, gold, platinum, iridium,

ım, rhodium, silver, and nickel.

Metals proper are by Berzelius divided into elecative or acidifiable metals, and electro-positive or ble metals. Under the former head he places selersenic, molybdenum, tungsten, antimony, tellurium, ium or tantalum, and titanium. Under the head of positive or basifiable metals, he places gold, platismium, iridium, palladium, silver, mercury, copper, 1, tin, lead, cadmium, zinc, nickel, cobalt, iron, manand uranium.

- I am under the impression that each of the folmetals, being, agreeably to the same authority, of forming with a halogen body the electroe ingredient in a double salt, should be considered lifiable; namely, gold, platinum, silver, palladium, , rhodium, uranium, chromium, titanium, molybdenanganese, osmium, mercury, nickel, copper, iron, c.
- When the objects which it may be desirable to are too numerous and complicated in proportion to e and attention which we have to bestow; we may such time as we have, either in a cursory, supernd indiscriminate examination of the whole, or in thorough study of the more important parts. Of courses I cannot conceive that any judicious peruld hesitate in choosing the latter.
- Lunder this impression I shall treat particularly of live metals proper, included in the following list—atinum, silver, mercury, copper, lead, tin, iron, zinc, ny, bismuth, and arsenic. Besides the metals thus ned, there are in the same class, palladium, rhodium, osmium, nickel, cadmium, chromium, cobalt, con, manganese, molybdenum, titanium, tungsten, n, and vanadium. Of these I shall give only a count, with descriptions and illustrations of their and useful properties, where such exist.

I subjoin a list of metallic radicals, comprising all the metals tellurium; which has been treated of as a basacigen element. car knowledge extends, the dates at which these metals severally mown, and the names of their discoverers, are mentioned.

Table of Metals classified as Metallic Radicals, also of the data & which they were discovered.

Names of Metals.	Authors of the discovery.							Dates of the Discovery.	
Gold		Known to the Ancients.							
Tin Antimony Bismuth	1	Described by Basil Valentine							1.000 1.530 1.015 couldry-
A reactive		Brandt							1733
Cobalt Platinum Nickel Manganese Tungeten Molybdenum	7	Wood, assay Cronstadt Gahn and S D'Elhuyart Hielm Klaproth Gregor Vauquelin Hatchett	y-Diag	ter, J	amai	ca.			1741 1761
Manganose .	1	Gahn and S	School	la					1974 1981
Molydenium	ij	Hielm .			*	-		•	1280
Dranium	1	Klaproth		:					1789
Tîtaniem	- 1	Gregor				4			1,3192
Titanium		Vauquelin		•					1797
Columbium . Palladium	- 5	Hatchett	*			1	4		3668
Rhodium	- 81	Wollaston		٠					1803
l ri dium	- ' [Descotils at							1803
Oamigza		Smithson T							1.806
Cerium Pot assium . .	1	Hisinger ar	id Bet	rselit	2.0		4		_
Bodium	- 11								
Barium	- 1	Davy .			4	P		4	1800
Strontium Calcium									
Cadmium	1	Stromeyer							1918
Lathium		Arfwedson	6		4			+	1818
Aluminium .	-)	WW							
Glucinium	- {	Wöhler	+				т		1826
Yttrum	7	Berzelius		_					1899
Magnesium .		Bussy .		-		•	4		1829
Vanadium .		Sefetrom	-						1830

Of the Generic Characteristics of the Metals.

1410. When newly cut, metals have a peculiar lustre. They are the best conductors of heat and electricity; the worst radiators and best reflectors of heat. All combine, directly or indirectly, with all the baseign bodies in one or more proportions. (633.) They are all susceptible of solidity and fluidity, and probably of the aëriform state. Mercury and arsenic are easily volatilized; and gold, silver, and platinum, though very difficult to burn or volatilize, are nevertheless dissipated by means of the compound blowpipe, galvanism, or electricity.

Of Properties possessed by some Metals, but not by others.

1411. The properties which come under this head, are permanency of lustre in the fire and air; malloability; ductility; elasticity; sensibility w

magnet; susceptibility of the welding process, and of acquiring, by a ion with carbon, silicon, or aluminium, the capability of hardening by ing suddenly refrigerated from a red-heat; also of being hardened by

> hammer, and of being restored by heat in the annealing process.

1412. The metals remarkable for permanency of lustre, are gold, platim, iridium, palladium, rhodium and nickel, called on that account noble, perfect. Those principally remarkable for malleability, are gold, silver, tinum, copper, palladium, nickel, iron, tin, cadmium, and lead. Among se, iron and platinum only, can be advantageously hammered at a very h temperature.

1413. The metals distinguished for elasticity, are iron, copper, and Iron, in the state of steel when duly tempered, is pre-eminent for

s property.

1414. The metals remarkable for ductility, are gold, iron, either pure, as steel, silver, copper, platinum, tin, and lead. In large rods or pipes, d and tin are the most ductile.

1415. The magnetic metals are iron, whether pure, in the state of steel, in that of protoxide, nickel, and cobalt. Those susceptible of the weldg process, are iron and platinum. Iron only is capable of uniting with rbon, silicon, or aluminium, and hardening consequently by quick regeration. Gold and platinum are distinguished by their superior gravity, hich is between two and a half, and three times as great as that of iron, 4 or zinc.

1416. All the metals have a specific gravity greater than five, if we ex-

pt those of the earths and alkalies.

1417. Of the Annealing Process.—Malleability, ductility, and toughses, in metals susceptible of the annealing process, are probably dependent n the quantity of caloric remaining in combination with their particles, hile in the solid state. When malleable metals are hammered, they give ut heat, and become harder, more rigid, elastic, and dense, until they acuire a certain maximum of density. This being attained, they are fracred if the hammering be carried further. Exposed to the fire until Mened, they are found on cooling to have regained the properties of rhich percussion had deprived them; and they may be again condensed, ented, and hardened, by the hammer.

1418. Of Alloys.—This name is given to the compounds formed by be union of different metals. There is always copper in gold and silver vin; and in the metal employed under those names by the smiths and wellers, there are various proportions of the baser metal. Brass consists f copper and zinc; pewter, of lead and tin, or of tin, copper, and anti-

lony.

If the Oxidability of Metals by Exposure to Air or Moisture, with or without Heat.

1419. Gold, silver, platinum, palladium and rhodium, do not become oxilized by exposure to water or oxygen at any temperature; and when oxilized by other means, on being ignited are reduced.*

The verb to reduce, has long been employed by chemists to signify the deoxiisomest of a metallic oxide, so as to effect its restoration to the metallic state, or int of a regular, to use another word which I shall also employ, to avoid circumlostion; although it is now somewhat antiquated. The verb to revive has been used the same sense as to reduce.

water alone, unless aided by a red-heat. Of these metals iron is made acted upon by the joint influence of air and moisture, at the ordinary to peratures of the atmosphere. Copper, tin, and lead do not decompate water at any temperature, but are oxidized at a red-heat, or at temperature sufficient for their fusion. Mercury is not oxidized by water under any circumstances. It is oxidized by agitation, or by a heat just below he boiling point, with access of air; but when distilled, it abandons the express which may have united with it previously.

OF METALS OF THE EARTHS PROPER.

1421. The metals included under this head are denium, glucinium, yttrium and thorium.

SECTION I.

OF ALUMINIUM.

1422. A chloride of aluminium was obtained by Oerstaby subjecting to a current of chlorine, an intimate mixture of alumina and carbon, heated in a porcelain tube. The affinity of the carbon for the oxygen of the earth, and of the chlorine for the metallic radical, was productive of carbonic oxide in the state of gas, and the chloride of aluminium in the state of vapour; of course the former escapes, while the latter condenses, within a glass tube purposely luted to that in which the materials are ignited, as already explained.

1423. By heating, with potassium, the chloride obtained by the process above mentioned, Wöhler liberated aluminium through the superior affinity of potassium for chloride.

rine.

1424. I have repeated this process so far as to obtain the chloride, and to expose it to reaction with potassium, but I found it difficult to extract the aluminium from the

residual mass to a satisfactory extent.

Wöhler obtained this metal, it is described as a gray powder much resembling that of platinum. Some little facets, which have sufficient magnitude to be distinguished, after compression under the burnisher, display a metallic brilliancy. Yet in the pulverulent form, the metal has so little power to conduct electricity, that, when interposed in the galvanic circuit, it interfupts the action. It is alleged, however, that, in a minute state of division, iron is a non-

tor of electricity, and Turner states that, by fusion, um becomes a conductor. It appears to me that its claims to the metallic character are not superior e of carbon in the form of plumbago. Its atomic is 14.

Aluminium burns with a heat so intense, as to he fusion of the resulting oxide, which becomes, on , hard enough to cut glass. Aluminium is not oxiden water is evaporated from it at a gentle heat. Siling heat it evolves hydrogen feebly, and the evolutionary once commenced, continues for some time frigeration. With concentrated nitric or sulphuric luminium has no reaction at ordinary temperatures; sisted by heat, it forms a sulphate or nitrate, acoxygen from one portion of the acid, and uniting e remainder. When subjected to a solution of potda, or ammonia, aluminium, by the decomposition water, is converted into alumina, which unites with ali, forming an aluminate. On this account, in prealuminium, there should not be an excess of potasand any potash produced during the process, should old the removed by the employment of a quantity of the larger than necessary.

Of Alumina.

. This earth is found nearly pure in the gems called ellers oriental, and classed by Brongniart, under the Corindon Telesie. The ruby, sapphire, amethyst, az, of the most beautiful kinds, are thus designated stony minerals they have the highest specific grand are only inferior to the diamond in hardness. In grown each other only in colour, they yield by anattle else than pure alumina. There are other jewels same name and colour, which ought not to be conlimited to the with those here alluded to. As an ingredient in hich owes its plasticity and all its striking qualities lina, this earth enters largely into the structure of estrial globe.

The spinelle ruby, a precious stone, and Gahnite, minntes,—the former of magnesia, the latter of which, however, there are six times as many atoms

ina, as of the other constituent.

one found in the United States, in the form of a stalant white and semitransparent, called Gibbsite; the other Siberia, called disapore, from the property of flying in pieces, or even powder, when heated, in consequence, doubt, of the vaporization of the combined water.

1430. Preparation.—Berzelius alleges that the alum commerce, if it contain oxide of iron, should be dissolt and recrystallized several times; or a solution being and allowed to stand for some time, the oxide of iron precipitated in yellow flocks. To the solution of alm a boiling heat, a solution of carbonate of potash is to added in excess, and the whole is to be digested at all derate temperature, to decompose any supersulphati alumina which the alkali may have precipitated. precipitate, after having been collected and well was upon a filter, is to be redissolved in chlorohydric a and precipitated by an excess of ammonia, either car or carbonated. This second precipitation is necessitation to get rid of a portion of carbonate of potash, with w the alumina forms a triple combination which cannot decomposed by water. The precipitate produced as mentioned, is to be collected and carefully washed. W dried it forms a hydrate, which, by a red heat, is t verted into pure alumina. One hundred parts of a yield a little more than ten of the earth.

1431. In France a species of alum is used, in whammonia takes the place occupied by potash in the comon alum. By heat, which expels the acid and all pure alumina may be extricated from this compound.

1432. Properties.—Alumina is white, plastic when a tened, soft to the touch, adherent to the tongue, inoderce insipid, and infusible in the furnace. It is the only which was fused before the compound blowpipe was vented. Its property of contracting and hardening heat, was noticed when on the subject of Wedgworpyrometer.

1433. It is remarkable that, although quite insoluble water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, this earth abstracts and retains a quantity of water, the earth abstracts and retains a quantity of water, the earth abstracts and retains a quantity of water, and the earth abstracts and retains a quantity of water and a property of the earth abstracts and a property of the earth abstracts and retains a quantity of water and a property of the earth abstracts and a

with moisture, which it slowly relinquishes in dry

There is a remarkable difference in the appearable hydrate of alumina as obtained by precipitation oncentrated, or a weak solution of alum. In the ase it is a white, friable, spongy powder, which is to the tongue, and, by exposure to a red-heat, h all its water. In the latter it forms a transparaw mass, which breaks by the heat of the hand mooth and conchoidal fracture, does not adhere ngue, or swell by the addition of water. In this e hydrate of alumina does not part with all its ren at a temperature above that of redness.

Alumina has a great affinity for vegetable colourers, which it consequently precipitates from their , forming the pigments known under the name of

This earth and its salts are of great use in dyenordants to cause the dyes to adhere. The latter, cases, have no affinity for the organic fibres which e dyed; but the alumina, combining with both the

the fibre, associates them permanently.

Alumina is soluble in solutions of caustic potash i, and even in those of baryta and strontia, but in liquid ammonia, only to a very small extent. has an affinity for oxybases so strong, as to be ed as acting the part of an acid in some instances. e acid and base of alkaline carbonates it forms mpounds, which will bear a low red-heat without the acid, or producing a more intimate union the earth and alkali.

The affinity of alumina for magnesia is so strong n separated simultaneously from a common solformer cannot be taken up entirely by the alkali, h the separation is effected. If magnesiferous after having experienced a red-heat, be subjected hydric acid, a white powder remains, which is an e of magnesia.

Three properties serve to detect alumina; first, its potash, and consequent solubility in a solution kali; secondly, the property which it has of form-sulphuric acid and potash, alum, so readily recogits crystallization and taste; thirdly, the property

of producing a fine blue colour, when moistened with nitrate of cobalt, and exposed to a strong heat.

exemplification of the commutable character of electrochemical characteristics. While with the alkalies and kaline earths it performs the part of an acid, with various acids it acts as a base, forming with them compounds both natural and artificial. Among the former is the mineral generally designated as feldspar, which is composed of silicate of alumina, and silicate of potash. Porce, lain is an artificial silicate of alumina. Its existence as the base in alum has been mentioned.

1441. Alumina was named from alumen, the Latin appellation for alum. The specific gravity of alumina is 2. It is composed of two atoms of aluminium, equivalent 23, and three atoms of oxygen, equivalent 24 = 52. It is, therefore, a sesquioxide.

Experimental Illustrations.

1442. Alumina, precipitated from a solution of alum by an alkali. Rendered blue by a solution of nitrate of cobalt. Contraction sustained by exposure to heat, illustrated.

Of Chloride of Aluminium.

1443. The chloride of aluminium is obtained as I have stated above. (1422.) It is partially translucid, lamellated in structure, of a greenish-yellow colour, and an astringent taste. Litmus is reddened by the action of this chloride. It dissolves in water with a hissing noise. When the solution is highly concentrated, it deposites crystals, which, being convertible by heat into alumina and chlorohydric acid, probably consist of one atom of chloride of aluminium, and one atom of water. According to Thenard, the chloride of aluminium forms, with the chlorides of potassium and sodium, compounds indecomposable by a red heat. These may be considered as formed by the union of a chloracid with a chlorobase.

SECTION II.

wo **a** ow

OF GLUCINIUM.

1444. Glucinium may be obtained from its oxide, glucina, by a process analogous to that above described for obtaining the radical of alumina. This metal resembles aluminum in appearance, and in many of its properties, but differs from it in not being susceptible of oxidizement by a solution of ammonia, or by boiling water.

Of Glucins.

Glecina is white and tasteless. It is insoluble in water, but forms with it a high is somewhat adhesive, but not sufficiently so to be moulded. It does not

y exposure to heat.

It is soluble in the caustic fixed alkalies, but not in ammonia. It likewise in the alkaline carbonates, and in that of ammonia especially, by which it ruished from alumina, as well as by its incapacity to produce alum, or to asblue colour when treated with nitrate of cobalt. It forms also a fluacid, rith the fluoride of potassium, precipitates from a hot solution in crystalline n the state of fluoglucinate of potassium (fluoride glucinico-potassique of s).

The equivalent of glucina is 26, being composed of one atom of glucinium,

at 18, and one atom of oxygen, equivalent 8.

Glucina exists in the emerald, comprehending the beryl and aquamarine: he suclass. In consequence of the peculiar sweetness of its salts, it was placina, from passes, sweet.

SECTION III.

OF YTTRIUM.

Yttrium was procured by a process quite analogous to that described for alult has a more metallic and crystalline aspect than that metal or glucinium. udes with oxygen and the acids are perfectly analogous to those of the mere mentioned. It is liable to be slowly oxidized in a solution of potash by emposition of water. Like glucinium, it is not oxidized by water even when

Of Yttria.

Yttria is insipid, infusible, and insoluble in water. It is uncertain whether ow tinge which it usually presents, is appropriate, or produced by impurities. dered snow-white by the presence of a small quantity of sulphuric acid. It is than baryta, being of a specific gravity approaching to 4.842. It is dised from other earths by its insolubility in caustic alkalies; while it dissolves carbonates, especially that of ammonia, although in a lesser quantity than

Yttria is principally characterized by its susceptibility of precipitation by rite of potassium (ferroprussiate of potash). Excepting thorina, this pro-

possessed by no other earth.

With acids it forms salts, having a sweet taste, and in some instances the the amethyst. In fact, the best means of detecting it, is the production phuric acid of crystals having this hue, which are extremely slow to dissolve, and which effloresce when heated. Its affinities are more feeble than those kalies or alkaline earths.

This earth has been found only in three Swedish minerals,—Gadolinite,

talite, and yttro-cerite.

Yttrium is composed of one atom of yttrium, equivalent 32, and one atom n, equivalent 8=40.

SECTION IV.

→*0 ● 0*****←

OF THORIUM.

Therium was first found, not many years since, in a single locality, in the wide or earth, combined with silicic acid. It is in the island of Loccun with, near the little village of Berwig, in Norway. It was found in I recembling obsidian, and called thorite, which contained 57 per cent. of waide of thorium, and in addition, lime, magnesia, iron, manganese, os-

mium, lead, tin, and a little alkalı combined with selicic acid and water. In m

the analysis of this mineral, Berzelius discovered therma

1456 From chloride of thorium, as from the other chlorides of the same genue, the radical may be evolved by means of potassium and heat. It may be extricated from the double fluoride of thorium and potassium, or fine. of potassium. Therium, in its appearance and in many of its properties, and sambles aluminum, but differs from it in not being exidized by reaction and ing water, dilute sulphuric acid, or alkaline solutions. When hexied goalfy l air, thonum inflames, and is converted into thorms.

1457 I do not conceive that either thorium, or any other of those substant merated as convertible by oxidizement into the earths proper, are more solidate

considered as metals, than carbon in the state of plumbago.

Of Thorna.

1458. Thorna is white, tasteless, and inodorous. In common with aluminars, and yttria, it is capable of acting as a base with water. The resulting to of thorna is by heat convertible into the anhydrous exide, in a state of great

1459. Thering may be known from its sulphate being more soluble in cold | hot water. It is composed of one atom of therium, equivalent 60, and one a oxygen, equivalent 8 == 68.

OF METALS OF THE ALKALINE EARTHS.

1460. Under this head are included magnesium, can barium, and strontium.

SECTION I.

OF MAGNESIUM.

1461. Magnesium was first obtained by Bussy in 1 by subjecting the chloride to the action of potassium, manner precisely similar to that already described for taining aluminium (1422). It resembles silver in co and fusibility. It is malleable, and has a decided me brilliancy. It is oxidized by exposure to the air, boiling water. When sufficiently heated in the air, it bines with oxygen, and is converted into magnesia. specific gravity is greater than that of water.

Of Magnesia.

1462. This earth exists abundantly in a state of bination in nature. Dr. Thomson states that a whole 1 of low hills, consisting of anhydrous carbonate of nesia, exist in India.

1463. Sulphate of magnesia is one of the salts v exist in the ocean, and, consequently, when sea wat evaporated in order to obtain common salt, the sulmay be obtained from the mother-water. For the n w of the salt first abovementioned, magnesia has been y obtained in this country, from an American mineral magnesite, which is silicate of magnesia, iron, and Many varieties of lime-stone and marble contain sia. The marble called dolomite, is especially well as a compound of lime and magnesia. The pre-of magnesia renders a carbonate less ready to give rbonic acid gas.

1. This earth may be precipitated from a solution of salt, by adding a solution of potash or soda. It kewise be obtained from the carbonate by heat.

5. Properties.—Magnesia is white, has a feeble alkaste, and affects vegetable colours like an alkali, feebly. (1070.) It is nearly insoluble in pure water, solves to a considerable extent in water containing ic acid, forming a soluble supercarbonate.

Magnesia is distinguished from the other alkaline not only by being less energetic in its affinities kaline properties, but by the solubility of its sul-

7. Magnesia is one of the most fixed and refractory nces in nature, and was deemed infusible until fused in 1801, with the aid of the compound blowpipe. pecific gravity of magnesia is 2.3, and its equiva-

Experimental Illustrations.

3. The precipitation of magnesia from a solution of salt, exhibited; also its effects upon vegetable

SECTION II.

CIUM, BARIUM, AND STRONTIUM, THE METALS OF E THREE PRE-EMINENTLY ALKALINE EARTHS.

that I deem it expedient to treat of them under one Their oxides constitute three of the four earths tished as alkaline, which are pre-eminent in alka-(1070.) Next to oxygen, silicon, and aluminium,

calcium is probably the most abundant element in the tion. Barium is comparatively a rare product, and a tium, as an ingredient in our globe, is still more spatistributed than barium. Neither exists excepting in a bination, and for the most part in the state of oxid union with an inorganic acid, especially carbonic and phuric acid.

Of the Evolution of Calcium, Barium, and Stronting

1470. In the last edition of this Compendium, it mentioned, upon the authority of some of the mode proved treatises of chemistry, that Davy had iso calcium. During the winter of 1838, being engage some efforts for obtaining the metal abovemention was induced to re-peruse the original lecture in which distinguished chemist above named described the rest

his attempts to isolate the metals in question.

1471. It should be known, that by Seebeck, and by zelius, and Pontin, amalgams had been obtained of cium, barium, and strontium. From the amalgams discovered, Davy undertook to distil the mercury; b frankly declared that he was in nowise certain the had succeeded in this object. In the case of calcium his "most successful" experiment, "the tube broke an metal took fire" before the process was completed. sequently to the date of these facts, as far as I have enabled to learn, neither Davy nor any other manipu has succeeded in making a less abortive experiment that in which he was most successful. This justifie idea, that there has been some inherent difficulty v could not be overcome by the means to which he reso Agreeably to my experience, the weight of sixty grain mercury, which is the quantity which he alleges hims have employed, cannot, by the most powerful appar be made to take up a sufficiency of calcium to lea perceptible quantity of this metal when the mercury is tilled from the aggregate. And I fully concur with] in the opinion, that the temperature requisite to mercury from an amalgam, either of calcium, of ba or of strontium, is higher than glass will bear.

To enable the reader to judge of the justice of my remarks respecting the advanced by Davy, I will here quote his own language.
"That to obtain a complete decomposition was extremely difficult, since a

1472. Having in my treatise on galvanism, or voltaic electricity, given an engraving and description of my apparatus, and an account of my process for the evolution of the metals in question, I shall here only quote a few words

respecting their properties as observed by me.

Any liquid containing it; and afterwards, with tests, are the appropriate proofs of its presence. They all ank in sulphuric acid; were all brittle and fixed; and, for sion, required at least a good red-heat. After being spt in naphtha, their effervescence with water was, on the rst immersion, much less active. Under such circumances they reacted, at first, more vivaciously with hydric ther than with water, or even chlorohydric acid; because these liquids a resinous covering, derived from the naphia, was not soluble, while to the ether it yielded readily."

SECTION III.

OF LIME, OR CALCIA, THE OXIDE OF CALCIUM.

1474. This oxide exists largely in nature in combination with carbonic acid, forming all the varieties of marble nd limestone. Some kinds of white marble, especially hat of Carrara, so celebrated on account of its employment in statuary, consist solely of this earth combined with water and carbonic acid, uncontaminated by any

d-heat was required, and that at a red-heat the bases of the earths acted upon the lass, and became oxygenated. When the tube was large in proportion to the quanty of amalgam, the vapour of naphtha furnished oxygen sufficient to destroy a ut of the bases; and when a small tube was employed, it was difficult to heat the utused as a retort sufficiently to drive the whole of the mercury from the base ithout raising too highly the temperature of the part serving for a receiver so as barst the tube." "When the quantity of amalgam was about fifty or sixty grains, found that the tube could not be conveniently less than one-sixth of an inch in umeter, and of the capacity of about half a cubic inch. In consequence of these ficulties, in a multitude of trials I had few successful results; and in no case uld I be absolutely certain that there was not a minute portion of mercury still in mbination with the metals of the earths."

In a subsequent paragraph the distinguished lecturer adds: "The metal from ne I have never been able to examine exposed to air or under naphtha. In the se in which I was enabled to distil the mercury from it to the greatest extent, the be unfortunately broke while warm, and at the same moment when the air enred, the metal, which had the colour of silver, took fire and burnt, with an intense

hite light, into quicklime."

^{*} See Nicholson's Journal, Vol. XXI. for 1808; or, Tilloch's Philosophical Magae, Vol. XXXIII.

other matter. Hence, if the acid and water be expelled by heat, the lime will remain in a state of purity. Oystershells yield very pure lime by heating them to incardescence.

a very high temperature, the matter constituting the imparities is prone to enter into intimate combination with the lime, impairing its causticity, and susceptibility of the slaking process. No doubt this arises from a diminute of affinity for water. The lime of shells is sometimes partially converted into a sulphide, by sulphur derived from the animal matter.

especially if covered, than in an open fire; and if the best be too sudden, the carbonate may be fused without the expulsion of all the acid, which is afterwards more tensciously retained. The extrication of the carbonic acid is promoted by a current of steam, or of any other aëriform fluid. But steam is preferable, as it is more easily procured, and cannot be productive of impurity. The rationale is, that homogeneous aëriform particles interfere with each other more than heterogeneous, which, agreeably to the Daltonian doctrine, to a certain extent oppose no resistance to reciprocal intermixture and penetration.

1477. After the first calcination, Berzelius recommends that the lime be slaked, and again calcined in an open

crucible.

1478. Properties.—The colour, taste, and smell of this earth, are well exemplified in the best kinds of lime used in building (sometimes called quicklime), which is, strictly speaking, oxide of calcium, isolated from the water and carbonic acid usually united with it as found in nature.

base, with water, acting as an acid. (826.) The water becoming, in consequence, consolidated, abandons its latent heat, or caloric of fluidity. Hence great sensible heat is excited, and when the mass undergoing the change is large, ignition occasionally ensues. The lime is by these changes rendered pulverulent, and is said to be slaked. The process is called slaking. The slaked lime thus produced, is by chemists called hydrate of lime. (826.) Quicklime is productive of heat, even when triturated with snow. 1480. Water takes up about 75sth of its weight of this

rth, forming lime-water. On this a pellicle is generated, on after exposure to the air, by the union of the lime th the carbonic acid, which always exists in the atmobere.

1481. In lime-water, some metallic oxides are soluble, secially those of lead and mercury. It follows, from the inition of acidity and basidity, that in the resulting commds, the oxides of the metals proper act as acids, while it of calcium acts the part of a base. (629, &c.) The operty which lime has of affecting vegetable colours, an alkali, has already been noticed. (1065, &c.) rough lime is precipitated from the aqueous solution, own as lime-water, by carbonic acid, yet an excess of is acid being supplied, the precipitate is re-dissolved. It in this way, no doubt, that the water in limestone counies becomes charged with this earth.

1482. The hardening of mortar is ascribed by Berzelius the affinity between the lime and the silicic acid in the

nd. Hence the necessity of this ingredient.

Experimental Illustrations.

1483. Characteristic changes produced in vegetable cours by the solution of the earth in water, called limeater. A glass of lime-water is not made turbid by air om a bellows, but becomes so on propelling the breath rough it. Absorption of carbonic acid by lime-water, hown. Solution of lime by an excess of the acid. Lime recipitated from solutions of its muriate or nitrate, by alphuric or oxalic acid.

Of Peroxide or Bioxide of Calcium.

1484. Oxygen is absorbed when passed over lime heated incandescence. By adding lime-water to oxygenated ater, acidulated with muriatic acid, Thenard procured ystals of bioxide of calcium. (853, &c.)

Of Baryta.

1485. This earth was named from the Greek Bugos, heavy; cause the minerals containing it are peculiarly heavy, hen compared with other earthy substances.

1486. Preparation.—To procure baryta, eight parts of the sulphate, finely pulverized, should be intimately mangled with one of charcoal, and afterwards triturated with two parts of resin, sugar, molasses, or wheat flour. The mixture is to be kept at a white heat, in a Hessian cruci-

ble, for three-quarters of an hour.

1487. The sulphate of baryta, by being deprived of oxygen, becomes converted into a sulphuret of barium, which yields a nitrate of baryta on the addition of nitric acid. The filtered solution by evaporation yields crystals of the nitrate, which should be decomposed in a porcelain or plastinum crucible. This operation is tedious; since the beat cannot be urged beyond a certain degree of intensity without causing the salt to rise up in a foam, so as to overflow the crucible. If the heat be arrested at a certain stage of the process, Berzelius alleges that a portion of a trous oxide remains united with the earth, forming a compound which has been mistaken for bioxide ("suroxide") of barium.

of strontia are, like those of lime, decomposable per se by heat. The addition of carbonaceous matter enables us to decompose them; as it changes the carbonic acid into carbonic oxide, which has no affinity for the earths, and,

therefore, escapes.

1489. Properties.—Baryta is acrid, slakes like lime, and is more soluble in water. It is more actively alkaline, both as respects its taste and its action on vegetable colours, than any other earth. It is gray at first, but absorbs water and becomes white. Its aqueous solution is rendered milky by carbonic acid, and, by combining with it, becomes covered with a pellicle of carbonate, when exposed to the atmosphere. From its solution in boiling water, baryta crystallizes on cooling.

1490. Solutions of barium, whether in the state of a hydrate, acetate, nitrate, or chloride, are very useful as tests for sulphuric acid, which, combining with the oxide of barium (baryta), previously existing in the hydrate or bitrate, or formed from the chloride by the decomposition of

water, is precipitated by them from any liquid.

1491. Ignited intensely, it absorbs oxygen if exposed to it, and passes to the state of bioxide. This earth is poison-

of one atom of barium = 69, and one of oxygen = 8 = 77.

Experimental Illustrations.

1492. Baryta, free from water, exhibited; also in crystals. Barytic water rendered milky by the carbonic acid of the breath. Solutions of baryta, and of sulphuric acid, introduced into distinct vessels of pure water, have no effect; but portions mingled in the same vessel produce cloud. Water, coloured by alkanet, turmeric, &c., thanged by baryta, as by an alkali.

Of Strontia.

1493. This earth is very analogous to baryta in its properties and composition. It is distinguished from baryta, by the red colour which its solutions communicate to flame, by its crystallization, and by its being more soluble in boiling water and less so in cold. Excepting baryta, it is more actively alkaline than any other earth, both as respects taste and its action on vegetable colours.

1494. Strontia water is not like that of baryta precipitated by a dilute solution of the sulphate of potash, or that of soda, and when added to a solution of bichromate

of lead its power as a precipitant is inferior.

1495. Strontia may be obtained from the carbonate or sulphate, by a process in every respect similar to that which has been described as the means of procuring baryta.

1496. The equivalent of this earth is 52.

Experimental Illustrations.

1497. Turmeric, alkanet, and red cabbage, changed by strontia-water, as by alkalies. Red colour of the flame of alcohol, containing strontia, shown. Effects of the aqueous solutions of the alkaline earths on a solution of bichromate of lead.

Of the Peroxides or Bioxides of Barium and Strontium.

1498. When the protoxides of barium and strontium

are heated in contact with oxygen gas, they absorb it, and are converted into bioxides. When an aqueous solutor of these earths is added to oxygenated water, the bioxides of their metallic radicals are precipitated in a crystalline form.

1499. It was by means of a bioxide of barium thus procured, that Thenard was enabled to obtain oxygenated water. (853.) The bioxide of barium was dissolved in chlorohydric acid. By adding sulphuric acid, sulphate baryta was precipitated, in which half of the oxygen of the bioxide was retained, the other half being left in combinate tion with the water of the solvent. This operation being repeated several times, the liquid became more and more surcharged with oxygen. Afterwards, the chlorine of the acid was precipitated by sulphate of silver, and the sale phuric acid, thus introduced, by baryta. Finally, the bid oxide being less susceptible of vaporization than water this liquid was removed by evaporation in vacuo over set phuric acid. (399.) Thus isolated, the oxygenated water was ascertained to deserve the appellation of bioxide, being found to hold two equivalents of oxygen for one of hydrogen.

OF THE METALS OF THE FIXED ALKALIES, OR ALKALIFI-ABLE METALS, POTASSIUM, SODIUM, AND LITHIUM.

SECTION I.

OF POTASSIUM.

1500. The discovery of potassium and sodium was made by Sir Humphry Davy, in 1807, by exposing their oxides, potash, and soda, to the divellent influence of the Voltaic current. These metals were afterwards obtained more copiously, by subjecting the alkalies, in contact with iron in a divided state, to intense heat in a luted gun barrel. Latterly, they have been obtained, with still greater facility, by heating the carbonates intensely, while intermingled with charcoal."

^{*} In Brunner's process, bitartrate of potash, or carbonized cream of tartar, which consists of carbonate of potash intimately intermingled with the residual carbon of the decomposed tartaric acid, is subjected to intense heat in a luted from mercury both, some coarsely powdered charcoal being added. The potassium was conveyed into a copper vessel containing naphtha as it came over from the bottle. For this result have substituted an iron tube, which becomes finally full of the metal and a carbon.

- 1. The alkaline metal, whether potassium or sodium, rolatile at any temperature above redness, is extriin the state of vapour, and condensed in a part of paratus where the heat is below redness.
- L. Properties.—Potassium, when newly cut, strongly les silver in appearance. It is malleable, and so ordinary temperatures, as to be moulded between gers like wax. When cooled to 32°, it becomes and exhibits, when broken, a crystalline fracture. s at 106°, and is converted into vapour when heated tle below redness. When exposed to the air at the ry temperature, it absorbs oxygen rapidly, and is ted into potash. This absorption is sometimes so especially when aided by friction, as to cause the nation of the potassium. I once lost half an ounce ussium, in consequence of attempting to extricate it ding the containing bottle by a file; it took fire, and tirely oxidized. The affinity of this metal for oxyso strong, that, when thrown upon water or ice, it es with the oxygen; while the hydrogen takes up a portion of the potassium, and burns with a beautie-coloured flame. Potassium is lighter than water, cific gravity being only 0.86. It is a good conducneat and electricity. Its atomic weight is 40.

SECTION II.

OF SODIUM.

1. Properties.—Sodium resembles potassium in its ance, and in many of its properties. It retains its

mass, which sublimes during the operation. The tube is then removed, and sarest the bottle screwed into a tapering tube, while the other orifice is a cap, into which it fastens by screwing. The tube is then placed vertically co, through the bottom of which the tapering tube extends so as to be out y of the heat. Under the orifice of this tube, a vessel may be placed conne naphtha, to receive the potassium as it descends in globules, after fusion sation from the state of vapour. The last portions are not evolved before the furnace reaches a white heat. The principal source of disappoint-brunner's process, is the failure of the luting. When this happens, the is soon burnt through. I have found it advantageous to secure the iron through in this process, while supported vertically in the furnace, by a stout fine same metal, the whole resting upon an iron disk supported by bricks assertant earth.

means, I procured last winter, at one operation, more than six ounces of

280 sodium.

softness and malleability when cooled to 32°. A globule of sodium, thrown upon water, swims to and fro on the surface with great rapidity, absorbing oxygen, and evolving hydrogen from the water; yet no inflammation ensues. This is probably owing to the rapidity of its motion, which, by bringing it in contact with successive portions of water, keeps the temperature below that which is necessary to combustion; since, when the water is thickened with a little gum, which tends to impede the motion of the globule, sodium burns with a brilliant yellow fiame. The presence of an acid produces the same result. The affinity of sodium for oxygen, does not appear to be so strong as that of potassium; since, according to Thenard, it is not oxidized when exposed to dry atmospheric air, or oxygen. It melts at 194°, and for volatilization, requires a higher temperature than potassium.

1504. Sodium forms a number of alloys with potassium; one of these remains fluid at 32°, and is lighter than naphtha. The specific gravity of sodium is 0.97223. It is a good conductor of heat and electricity. Its atomic

weight is 24.

Experimental Illustrations.

1505. The inflammation of potassium and sodium upon water and ice, exhibited; also the regeneration of the alkali, demonstrated by the usual tests. The decomposition of potash, by iron card-teeth, heated to incandescence. Apparatus for its evolution, exhibited.

Of Potash or Potassa, and Soda, the Protoxides of Potassium and Sodium.

1506. A ley, obtained by the lixiviation of the ashes of inland plants, especially wood, when boiled down, yields the potash of commerce. Potashes ignited so as to destroy vegetable colouring matter and other impurities, again dissolved, and boiled to dryness, form pearlash. Pearlash dissolved in water, boiled with lime to remove the carbonic acid, filtered, and boiled down to the consistency of moist sugar, dissolved in alcohol, and boiled down gradually, and lastly, fused at a red-heat in a silver vessel, forms the potash, or, more strictly, the hydrate of potash of chemists. If, as soon as the alcohol has escaped, the residual mass

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be refrigerated, it crystallizes. After fusion at a red-heat, the alkali contains about 20 per cent. of water, existing in it as an acid, and of which, per se, it cannot be deprived by heat.

1507. Pure carbonate of potash may be procured from bitartrate of potash, whether carbonized by heat, or deflagrated with pure nitre, by subjecting the residue to water, and the resulting solution to heat, to vaporize the solvent.

1508. To obtain pure potash, or in other words, to remove carbonic acid from the alkali of a carbonate, Berzelius advises the addition of one and a half parts, by weight, of pure hydrate of lime, to one part of a pure carbonate, obtained as abovementioned, dissolved in a cauldron, and kept boiling. The lime is not to be added at once, but gradually; as without this precaution, the resulting carbonate of lime retains, like a sponge, a great part of the alkali. The liquid is to be tested by an acid or by lime-water, antil it ceases to indicate the presence of carbonic acid. After this, it may either be kept in a liquid state, or evaporated till it crystallizes, and preserved in crystals; or being ignited till it becomes fused, and poured out on a slab, or into moulds, it may be preserved in the state of hydrate.

1509. I have used for the purpose last mentioned, the moulds usually employed for casting musket balls. The spherical form presenting the least surface in proportion to the mass, is favourable to the preservation of a substance liable to be deteriorated by contact with the atmos-

phere.

1510. The crystals of potash, thus procured, are always free from carbonic acid, and if derived from a pure carbonate, excepting water, may constitute pure potash. But when pearlash is the carbonate employed, alcohol must be resorted to, after the caustic ley has been evaporated to the consistency of moist sugar, in order to get rid of the impurities. This liquid combines with the pure potash, while a portion of water contained in, or formed from, the alcohol, separates from it in union with the impurities.

Vhich grow on the sea-shore, as potash is by the incineration of those which grow inland. It is procured also from

chloride of sodium, and sulphate of soda.

1512. Soda is purified, and procured in the state of hy-

drate, or in crystals, by a process analogous to that above described for its kindred alkali.

1513. Properties of Potash and Soda.—Potash and soda are of a grayish-white colour, and, in common with other alkalies, have a peculiar taste. They render tincture of turmeric brown, syrup of violets green, and alkanet blue. Colours changed by acids, are restored by them. They are the opposites of, and antidotes to, acids, and capable of forming with them neutral compounds, or, in other words, such as are neither acid or alkaline. They are incorrectly said to render vegetable blues green, as if this were universally true. Alkanet is made blue by them, while neither litmus nor indigo is rendered green. (1065, &c.)

1514. Although potash is more soluble than soda, and is deliquescent, while soda effloresces; yet the salts of soda are more soluble than those of potash. Both cauterize the flesh. Potash is the more active. Common

caustic is an impure hydrate of this alkali.

1515. Crystallized potash contains four atoms of water to one of the oxide, of which three only can be expelled by heat. After fusion it may be called, however paradoxical it may seem, an anhydrous hydrate, though not an anhydrous oxide. Both potash and soda fuse when subjected to a red-heat. The atomic weight of potash is 48, that of soda, 32. The hydrate of potash consists of one atom of alkali, and one of water.

1516. Potash may be distinguished from soda, by its forming salts nearly insoluble in water with tartaric, or oxychloric acid; while those formed by soda with the same acids are soluble. Chloroplatinic acid causes a yellow precipitate with potash, but not with soda.

Experimental Illustrations.

1517. Characteristic changes produced in vegetable infusions, as in previous illustrations. (1075.)

1518. To a saturated solution of potash and soda, or their carbonates, a saturated solution of tartaric acid being added in excess, crystals are yielded by the potash only. Into different salts of the two alkalies in solution, chioroplatinic acid being poured, a yellow precipitate distinguishes the potash.

283 sodium.

Of the Peroxides and Suboxides of Potassium and Sodium.

519. Peroxide of potassium is produced by the combustion of potassium on a te of silver in oxygen gas, in which case the metal acquires three times as much ygen as it holds in the state of potash. The peroxide is also obtained when nitre intensely heated, or when potassium is deflagrated with nitre.

1520. Two parts of sulphate of potash, ignited intensely with one of lampblack,

e a pyrophorus which takes fire spontaneously with scintillations in the air. This ses, no doubt, from the extreme state of division in which carbon, potassium, and

phur exist in the residual mass.

1521. The peroxide of potassium is of a greenish-yellow colour, and possesses st of the properties of the protoxide, excepting that of acting as a base. When night in contact with water or acids, it is decomposed into potash and oxygen.

1522. The peroxide of sodium is of a greenish-yellow colour also, and, in its prorties, is analogous to the peroxide of potassium, except that at a high temperature ibandons part of its oxygen, and is converted into protoxide. It cannot, theree, be obtained by burning sodium in an excess of oxygen; since the heat produced the combustion, would decompose the peroxide, if already formed. In order to cure it, it is necessary to heat soda in contact with oxygen. The peroxide of so-

am contains one and a half atoms of oxygen, united to one of metal.

1523. Berzelius mentions that suboxide of potassium may be obtained by heating e metal in a quantity of oxygen inadequate for its saturation; also by exposing to a nperature of about 40° F., a mixture of hydrate of potash and potassium, in equilent proportions; in which case the metal is oxidized at the expense of the comed water, the hydrogen escaping. The anhydrous protoxide may be obtained in manner, by heating potassium with a greater quantity of the hydrate. Turner ges, however, that the suboxide of potassium is generally regarded by chemists nothing more than a mixture of potassium and potash.

524. According to Berzelius, a suboxide of sodium may be obtained by the same uns as the suboxide of potassium, substituting the one metal for the other. The e uncertainty, however, prevails with regard to it, as with regard to the suboxide

525. When potassium or sodium is heated in ammonia, it combines with nitrogen I liberates hydrogen, and the resulting nituret absorbs ammonia; so that we have ombination of two binary compounds of nitrogen, which may possess, to a small ent, the relation of acid and base. There are, however, no phenomena in chestry which are more anomalous than those which are associated with the producand evolution of this compound. Nevertheless, as its nature is unintelligible n to adepts, I shall not present the details here.

526. I hinted, when entering upon the subject of nitrogen, that it would be seen he sequel, that it was not destitute of pretensions to a place in the basacigen is. It was in reference to the phenomena above alluded to, that I made that re-

527. If nitrogen form the common ingredient in two compounds, one electrorative, the other electro-positive, which combine to form a third, it fulfils the conion of a body producing both an acid and a base, and is of course a basacigen ly. Yet it has already been pointed out that there is no class, of which some of members do not display properties which might cause them to be placed in anor class.

Of Phosphurct of Potassium.

528. Phosphorus and potassium, heated together in nitrogen or hydrogen gas, abine with the phenomena of combustion. In phosphuretted hydrogen, potassium ns, combining with phosphorus, and liberating the hydrogen. 529. This r losphuret decomposes water, but, according to Berzelius, the gas lived does not inflame spontaneously.

Of the Compounds of Potassium with Carbon, Boron, and Silicon.

1530. The black matter which remains after the distillation of potassium, as obby Brunner's process, is alleged by Berzelius to be a percarburet of potassium. hen moistened it inflames, no doubt by decomposing water, and evolving potassuted hydrogen. The black matter which obstructs the tube used in the evolution **potassium by** the process above mentioned, is also held to be a carburet. (1501.) 1531. These carburets I have found useful in purifying naphtha, by its distillation them. After undergoing this ordeal, potassium may be kept in it with less appenrance of reaction. I am under the impression that the carbon which remains in the iron bottle, is imbued with potassium, possibly in a state of chemical cause. This may be used likewise for the purification of naphtha

1532 It appears that, during the reduction of borie acid by potamium, a bornest is

formed, since a portion of the mass evolves a gas on being montened, which has not the smell of pure hydrogen. It is probably bornretted by drogen 1533. A siliciaret of potassium is obtained during the decomposition of flumber acid gas. A portion of the liberated silicon, combining with potassium, forms the compound in question. This, on being moistened, gives off hydrogen, which the a peculiar odour resembling that of phosphuretted hydrogen. The analogy between these results and those mentioned in reference to boron, is obvious.

SECTION III.

OF LITHIUM.

1534. A fixed alkali was discovered, in 1816, by Mr. Arfwedson, to cust in small proportion, as an ingredient in a mineral called petalite. He ofterwards discovered it in two other minerals, called spedumene and lepidelite. Allusion to this alkali, and the origin of its name, was made under the head of Ammonia. (1081.)

1535. By the influence of the Voltaic pile, decided indications have been obtained of the existence, in lithia, of a metallic radical. To this the name of lithium has been given. Lithium resembles sodium in appearance. It

atomic weight is 6.

Of Lithia.

1536. Lithia, known only in the state of hydrate, is white, caustic, and has all the attributes of an ulkali. When lithia, whether in the state of carbonate or uncombined, is heated in contact with platinum, the metal is attacked, and a compound is formed, which, according to Thenard, probably consists of the oxide of platinum, united to the oxide of lithium, and must of course be a platinate of lithia. Lithia is composed of one atom of lithium, equivalent θ_1 and one atom of oxygen, equivalent $\theta_1 = 14$.

1587. Lithia is less soluble in water or alcohol than soda or potash. It carbonate is less soluble in water than the carbonates of those alkalis-The chloride of lithium is deliquescent, and soluble in alcohol, the phate of lithia is insoluble in water; in which respects these companies differ from the corresponding combinations, formed by the other fixed alta-

lies, or their radicals.

Of the Reaction of Chlorine, Bromine, Iodine, Fluorist, and Cyanogen, with the Metals of the Earths and Alko-

1538. In a former edition of this work, it was mentioned that for aluminium, glucinium, yttrium, thorium, and magnesium, chlorine has not sufficient affinity to expel the oxygen from their oxides; and that it was only in the state of oxide that they could be subjected to the gas-

as been already stated, that Oersted ingeniously con-ad to enable chlorine to combine with aluminium, by co-operating affinity of intermingled carbon for the gen with which, in the state of earth, this metal is ed: also, that a similar process had been successfully loyed to obtain the chlorides of glucinium, yttrium, ium, and magnesium. The most important considera-, associated with the existence of these chlorides, is susceptibility of decomposition by potassium, and the equent isolation of their metallic radicals.

539. When the oxides of calcium, barium, strontium, ssium, sodium, and lithium are heated in chlorine, these als are converted into chlorides, the oxygen being dised. Potassium and sodium burn actively in chlorine, it appears probable that any of the metals of the als or alkaline earths may, with heat, if not without, be tly combined with any of the halogen bodies. e combinations may be obtained in the wet way by plex affinity, on presenting their oxides to the acids ied by these bodies with hydrogen.

540. The chlorides of the metals of the alkalies, and he alkaline earths, are all soluble, and some of them quescent. When in solution, they contain the same nents as if they were chlorohydrates of oxybases; and

therefore, considered as such by some chemists.

541. The difference between a chloride in solution and la chlorohydrate, is rendered evident by setting down ingredients agreeably to both suppositions, as fol-**5:**—

orine, hydrogen.	Oxygen, metal.
lorohydric acid.	Oxide.
Chlorohydra	ate.
ygen, hydrogen.	Chlorine, metal.
Water.	Chloride.
Dissolved	Chloride.

1542. The soluble chlorides produce white precipitates in solutions of silver, lead, or black oxide of mercury-They do not deflagrate with charcoal, nor do they, hke sulphates, after being heated with it, yield the odour of sulphuretted hydrogen on being moistened-

1543. The soluble chlorides of the metals of the alkaline earths and alkalies, excepting that of magnesium, arc, by heat, converted into anhydrous chlorides, easily detected

by the fumes which they give with sulphuric acid-

1544. Bromine, like chlorine, when heated with any of the fixed alkalies, or alkaline earths, except magnesia, displaces the oxygen and combines with the metallic radical Like chloring also, it does not, per se, produce this effect

either with magnesia or the earths proper.

1545. The affinities of iodine arc, in most cases, less energetic than those of chlorine or bromine. Potash and soda are the only oxides of the metals of the earths and alkalies, from which iodine can, with the assistance of heat, expel the oxygen, in order to combine with their metals.

1546. The bromides and iodides, when combined with water, may, like the chlorides, be regarded either as in a state of solution, or as bromohydrates and iodohydrates. The bromides may be recognised by the red vapours which arise, when they are heated in a tube with the bisulphate of potash. Similar vapours would be given out by the nitrites, if treated in the same way; but the bromides may be distinguished from those salts, by their not deflagrating when thrown on incandescent coals.

1547. An iodide may be detected by dropping a portion into sulphuric acid, heated nearly to the point of ebuilities. (738.) By these means iodine, if present, will be made apparent in the form of a violet vapour. Indine is also displaced from its combinations by chlorine; and, when these, previously to the addition of chlorme, are mingled with a paste made of starch, a blue colour is produced. It is alleged that sea sait sometimes contains a quantity of

iodine adequate to produce this result.

1548. Berzelius states that, when potassium is heated in cyanogen, it is converted into a cyanide; also that the habitudes of sodium are in this respect similar. It is probable that the same result would ensue with all the metals

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alkalies and alkaline earths. Cyanogen is usually ted by the reaction of potash with animal matter, leoxidizes the alkali, and at the same time furnishes he elements of cyanogen, which, in consequence, neously unite with each other and with the metal,

r a cyanide of potassium.

Nhen the cyanoferrite of potassium (ferropruspotash) is intensely heated, the cyanoferric acid is cosed. The cyanide of potassium remains mingled carburet of iron, and may be extricated by solution, on, evaporation, and crystallization. Subjecting the errite of sodium to a similar process, the cyanide of may be obtained. (1299, &c.)

In the cyanides may be detected by their power of ing a blue colour with solutions of the peroxide of lso by evolving the odour of peach blossoms, when

ed to chlorohydric acid.

. It is highly probable that the reaction of fluorine e metals of the earths and alkalies, will prove to be ous to that of chlorine. The fluorides, however, nuch from the chlorides in solubility. Some variethe fluoride of calcium constitute Derbyshire spar,

he chloride of calcium is a deliquescent salt.

The presence of fluorine in a mineral may, in a y of instances, be detected by the following process. mall portion of it be pulverized, and subjected to the about twice its weight of concentrated sulphuric a leaden, silver, or platinum cup. Let this cup be I by a glass plate, coated with beeswax, through some letters have been traced so as to denude the s surface. After exposure for half an hour, aided auch heat as can be used without melting the wax, is should be relieved from its coating and examined f the portions of the vitreous surface, exposed to les, prove to be so corroded as to render the chatraced through the wax distinguishable, the pref fluorine may be inferred.

. Berzelius informs us that when this principle is in ation with silicon, it will not act on glass. Hence ses that the mineral, suspected of containing fluocid, should be subjected to the flame of the blowcone end of a glass tube, of which both ends are that the fumes produced may be impelled by the

blast through the tube from one orifice towards the other. By these means, milky spots will appear on the glass in consequence of the condensation of water containing fluosilicic acid, if this be an ingredient in the mineral.

Of the Reaction of Sulphur, Selenium, and Tellurium, with the Metals of the Earths and Alkalies.

1554. Sulphur unites with all the metals of the alkalus and alkaline earths, so far as the experiment has been tried, whether presented to them in the metallic state, or in that of oxide. Its power of reducing their oxides it greater than that of any other basacigen body; as when present in excess, it acts by its affinity for the oxygen and the metal. (523, &c.) The affinity of the halogen bodies for oxygen, is so inferior to that of sulphur, that when oxygen is expelled from oxides by one portion of them, it does not combine with another, however great the excess

in which they may be present.

1555. Sulphides (sulphurets) are also formed by deoxidizing the sulphates by carbon or hydrogen with the aid of heat, (1436, 1437,) by boiling in water equivalent weights of sulphur and the earth or alkali to be combined; or by passing sulphydric acid into water, holding the oxide in solution or suspension. When this is done under favourable circumstances, the metal is converted into a sulphobase by the sulphur of one portion of the acid; while the compound thus formed unites with another portion of the acid, forming a sulphosalt, denominated a sulphydrate. This view of the subject we owe to Berzelus. who has shown that sulphur, selenium, and tellurium, all have the property of forming acids with one set of radicals, and bases with another; and that the sulphacids and sulphobases thus formed, are capable, like oxacids and oxybases, of forming compounds which he considers as sulphosalts, or salts in which sulphur performs a part analogous to that which oxygen performs in oxysalts, such as the sulphate or nitrate of potash.

an alkaline solution, sulphydric acid (sulphuretted hydrogen) combined with the oxybase, forming what was called a hydrosulphuret. It was also supposed that a sulphide of an alkalifiable metal, by solution in water, would be converted into an oxybase by the oxygen of the water; while

the hydrogen, with a double proportion of sulphur, forming bisulphuretted hydrogen, would combine with the oxybase.

1557. Through the sagacity and industry of Berzelius, much knowledge has of late years been acquired respecting the combinations of sulphur with the alkaline metals. He mentions seven compounds, in which, supposing the quantity of the potassium in each to be the same, the quantities of the sulphur are severally 1, 2, 3, 3½, 4, 4½, 5.

1558. To remember the details respecting the preparation and characteristics of these sulphides, would be too great a burthen for the memory of those who are not so

situated as to take a particular interest in them.

1559. Sulphides of the metals of the earths and alkalies, on being moistened with water, evolve sulphydric acid, and produce this result still more actively on being subjected

to chlorohydric acid.

- 1560. The selenides of the metals of the earths and alkalies may, in most cases, be produced by heating the metal with selenium. The selenides of these metals bear a great resemblance to the sulphides, and when heated are reduced to the metallic state, producing the smell of horse-radish.
- 1561. The tellurides are but little known, and, except so far as they act as telluracids or telluribases, so as to give pretensions to tellurium to be placed among the basacigen elements, they are uninteresting.

Experimental Illustrations.

1562. Sulphides in solution exhibited. Earths precipitated by acids.

OF METALS PROPER.

1563. The metals included under this head, are gold, platinum, silver, mercury, copper, lead, tin, bismuth, iron, zinc, arsenic, antimony, palladium, rhodium, iridium, osmium, nickel, cadmium, chromium, cobalt, columbium, manganese, molybdenum, titanium, tungsten, uranium, cerium, and vanadium.

SECTION I.

OF GOLD.

1564. Gold is usually found in nature nearly pure. It is not liable, the other metals, to be disguised by a union with oxygen or sulphur. The precipitate obtained from a solution of gold coin in aqua regia, by the green sulphate of iron, is pure gold. This metal is also purified by expanse to heat and air, or by natice acid, by which means baser metals are oxidead; as in the processes of cupellation and parting.

1565. From the sands, or ores, in which they exist naturally, minute portions of gold are collected by trituration with mercury, with which they amalgamate. The mercury is distilled from the amalgam thus obtained,

by means of an iron alembic.

1566. Properties.—The specific gravity of gold is 19.3, and its equation 200. Its colour and lustre are too well known to need description. It is the most malicable and ductile metal, and suffers the least by exposure to in and moisture. Gold leaf, which is about 1000 times thinner than printing paper, retains its lustre in the air. Gold leaf transmits a greenish light, but it is questionable whether it be truly translucent. Placed on glass, and viewed by transmitted light, it appears like a retina. It is erroceously spoken of as a continuous superfices.

1567. Gold fuses at a low white-heat, but requires the temperature produced by the compound blowpipe, by galvanism, or by the explosive power of electricity, to volatilize or oxidize it. Its not being hable to tarmsh by exposure, is due to the weakness of its affinity for oxygen or sulphur.

1568. When a solution of chloride of gold is mixed with sulphinic ether, the ether takes the metal from the chlorine, retaining it in solution. If iron or steel be moistened with this ethereal liquid, it is productive of a slight gilding.

1569. Phosphorus, carbon, and the baser metals, also hydrogen gas and its compounds, by a superior affinity for oxygen or chlorine, preseptate

gold from the solution of its chloride in the metallic form.

hydrogen of water. The hydrogen thus liberated, takes chlorine from gold, forming of course chlorohydric acid, which has no attenty for the metal, unless in the state of chloride. As oxygen is necessary to the bar of an oxysalt, so chlorine is indispensable to the constitution of a chlorous.

1571. The union of gold with mercury, was adduced as an exemption-tion of simple chemical combination. (515.) The compound thus formed, when the ingredients are in due proportion, is of great use as the mean of that kind of gilding which is the most firm and durable. The affacts between the mercury and copper, renders it easy to coat with the analysis the surface of any mass formed of this metal. Subsequently, the mercury may be driven off by heat, leaving a pellicle of gold upon the cupreous surface, which only requires burnishing, in order to display the colour and brilliancy of gold.

1572. With arsenic, gold combines energetically, absorbing this metal in the form of vapour, at a red-heat, without changing colour. Gold loss its malicability by acquiring \$\frac{1}{2}\dagger the of its weight of arsenic. Probably gold may be united with all the metals. Phosphorus forms with it a britise

compound.

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The affinity between chlorine and gold is pre-eminently energetic. ination ensues, whether the metal be heated in the gas, or presented iqueous solution, or in aqua regia, which is essentially a solution of in water. Aqua regia is produced by the mixture of chlorohydric ric acid. It ought not, however, to be considered as a combination

As soon as the mixture is effected, a decomposition of both of the mmences.

One atom of nitric acid, by yielding three out of its five atoms of (957,) can take all the hydrogen from three atoms of chlorine. (874.) se, three atoms of chlorine and one of nitric oxide are emancipated. cids employed be concentrated, both the nitric oxide and the chloevolved; but if there be a sufficiency of water, the chlorine remains with it, forming a more concentrated aqueous solution of chlorine n otherwise be made. Excepting that it contains chlorine in a legree of concentration, which of course enables it to act with more aqua regia does not differ, in its solvent powers, from a solution of in water. It cannot properly be considered as a distinct acid; only acts by imparting chlorine, being incapable, as an aggregate, ing into combination.

The name of aqua regia, or royal water, was given to this solvent, unt of its property of dissolving gold, the alleged king of metals. The promulgation of the French nomenclature, it has been called triatic acid; but as this conveys a false idea of its nature, I would by its old name, aqua regia, or, if a new name be necessary, I suggest that of nitrohydrous chlorine. Latterly Gay-Lussac has that iodic acid is a solvent of gold; and by Mitscherlich, the same ascribed to selenic acid. When boiled with three parts of sulphur, of potash, one part of gold is dissolved as an ingredient in a soluhosalt. (1541.)

Of the Compounds of Gold with Oxygen.

By subjecting a protochloride of gold to a solution of caustic potash, oxide um, the chlorine and oxygen exchange places; so that a protoxide of gold, ride of potassium result. The trioxide of gold is obtained by digesting an solution of the bichloride with magnesia in slight excess. This oxide, which e of acting both as an oxacid and as an oxybase, in this instance acting in a capacity, combines with the magnesia, and constitutes an aurate, of which ter part precipitates, while the remainder continues in solution. The precipulate washed with water until it ceases to acquire a yellow colour by the of chlorohydric acid. It should then be digested with nitric acid, which with the magnesia, and thus isolates the trioxide. If the nitric acid emeron concentrated, we obtain the trioxide in an anhydrous state, and of a brown but if dilute, as a yellowish-red hydrate.

The protoxide consists of one atom of gold and one of oxygen, the trioxide, f gold and three of oxygen. Hence, agreeably to the example of Thenard, ite it as a trioxide. (756.) Acting as a base, this oxide combines with nitric iric acid. It is precipitated from these combinations by water, which acts,

, in this case, as an oxybase of hydrogen. (826.)

As an oxacid, trioxide of gold unites with all the alkalies and alkaline earths. te of ammonia, a compound which explodes by percussion, has long been nder the name of fulminating gold. Berzelius alleges that there are two ie, containing an excess of ammonia, detonates more powerfully; the other, ith a lesser quantity of the alkali, contains chloride of gold, by which its enfeebled.

A precipitate, of a beautiful purple colour, may be obtained either by mixing stations of the chlorides of tin and gold; or by immersing an ingot of tin, I, in a solution of chloride of gold, containing some free chlorohydric acid. recipitate the name of purple powder of Cassius has been given. I infer account of this compound, given by Berzelius, that it consists of gold, tin,

hydrogen, and oxygen. Respecting the mode of combination there to come obscurity.

1580 In consequence of this property of producing the purple of Cassaus, im, whether in the metallic state or that of descrived protochloride, is the best test for

1581. Berzehus done not consider the purple powder into which gold is reduced by successive electric discharges, as any thing more than metallic gold in a state of minute division.

Of the Compounds of Gold with the Hulogen Bodies

1582. The protochloride of gold is obtained by exposing the trichloride to a grate heat, which drives off two atoms of chlorine, leaving the gold in combination with the remainder. If the heat be carried too far, it is upt to decompose the protochloride ride into metallic gold and chlorine. On this account it is better to stop the symmotion before the trichloride is entirely decomposed, and to wash the resulting mass with water, which removes the trichloride, and leaves the protochloride which is insolable in that fluid when cold. A solution of the trichloride of gold is obtained when gold is dissolved in aqua regis, any excess of chlorohydric and is expelled by heat. It is of a pale yellow colour, and has an astringent and disagreeable two. This chloride combines as an acid with the chlorides of the alkaline metals, forming thioroaurates. Hence I consider this as entitled to the appellation of chlorianing acid. The trichloride of gold, as its name implies, is composed of one atom of gold, and three atoms of chlorine.

1583. Bromine forms with gold a tribromide, which corresponds in composition and chemical properties with the trichloride of the same metal. The indule of gold agrees, in composition and chemical relations, with the protochloride of gold the cyanide of gold appears to act as an acid.

Of the Compounds of Gold with Sulpher.

1584 Gold forms with sulphur a protosulphide and a trisniphide. The protosulphide is formed by passing a current of sulphydric acid gas through a botting solution of the trichloride. It is of a deep brown colour, and is decomposed by heat mismetallic gold and sulphur. The trisulphide may be precipitated by passing a current of sulphydric acid into a dilute solution of the trichloride, or by adding an acid to a solution of the sulphurate of potassium. The trisulphide is of a deep yellow colour, and is decomposed by heat. With sulphobases it acts as an acid, but with the more powerful sulphacids as a base.

Experimental Illustrations.

1585. Some gold leaf is placed in two glass vessels. Nitric acid being poured into one, and chlorohydric acid into the other, the gold is not acted upon; but when the contents of the two vessels are united, the gold disappears.

1586. Gold, dissolved by aqua regia, and precipitated by sulphate of iron, or by chloride of tin. A cylinder of phosphorus, immersed in a solution of the metal, acquires the appearance of a cylinder of gold. Separation of gold from its solution by ether. Effects of the ethereal solution exhibited. Action of mercury on gold leaf.

SECTION II.

OF PLATINUM.

This metal is found in South America, and in Russia, in an imrular form, known as the native grains of platinum. In addition etal, the native grains contain several other metallic substances in combination or mixture. The aggregate thus described is, for the t, soluble in aqua regia; the habitudes of platinum, in this respect, s in others, being more analogous to those of gold than any other nature. On adding to a solution of the native grains of platinum, in ria, a solution of sal-ammoniac, an orange-yellow precipitates, but ible in water, is obtained. This being carefully washed and desicid finally exposed to a red-heat, in a platina, porcelain, or black rible, the metal is isolated in a mass so porous, as to have received of platina sponge, from its resemblance in structure to the well ubstance to which this name belongs. By extreme mechanical the platina sponge is so far consolidated that by intense heat and ng it is welded into a perfectly tenacious mass, having, in a high ll the attributes of a noble metal. (1404.)

I have lately been enabled, by an improvement in my hydro-oxy-pipe, to fuse twenty-five ounces of platinum into a malleable mass, at thus obtained, is less liable to flaws than that produced by the process above described. My process is especially important as us to unite old platina ware, or clippings, into malleable masses of nt dimensions, without re-solution in aqua regia. The necessity of is last mentioned course, reduces platina in that state, to a value

than 1 higher than that of the native grains. (394.)

According to Berzelius, platinum, as obtained by the process ntioned, is alloyed with iridium, and inferior to the pure metal in filliancy, ductility, and malleability; while at the same time it is and more suitable for the purposes for which it is usually emtrementation by chloride of potassium, igniting the precipitate, ng it, and precipitating again by sal-ammoniac; and lastly, by rese precipitate by ignition to the spongy form, from which by presthe welding process, it may be made coherent and malleable, as prementioned process for obtaining the metal.

Properties.—The colour of this metal, as ordinarily obtained, is ate between that of silver and steel; but when pure, as above resembles silver both in colour and softness, more than when with iridium. Its specific gravity is 21.53. A cubic inch of it ore than three-fourths of a pound. It is nearly twice as heavy as lead, heaviest body known. It is less ductile and malleable than gold, r and more tenacious; though, in these respects, inferior to iron, it is susceptible of being hammered and welded at a white-heat. There is exidized nor melted by the highest temperatures of the airmorge. It was first fused in a focus of the solar rays, afterwards of a stream of oxygen gas on ignited charcoal, but much more the compound blowpipe, under which it was first oxidized and dispheat. It fuses and burns easily in the Voltaic circuit, and is

dispersed and oxidized by mechanical electricity. It is one of the west

conductors of heat among metals.

1591. In its habitudes with oxygen, chlorine, and the acids, it is analogous to gold, being, like that metal, detected by protochloride of tim, which produces with it a claret colour. It unites so energetically with tip at red-heat, as to occasion the phenomena of combustion. (348.) When is a divided state, as obtained by igniting the chloroplatinate of aminonium,

amaigamates with mercury by trituration.

1592. Platinum combines with boron, silicon, and phosphorus. On account of its infusibility at the highest temperatures produced by the archivenace, or forge, and its insusceptibility of being corroded by the acids usually employed in chemical processes, it is much used by chemists for cracible evaporating vessels, and spoons; also in experiments in which Voltage cracing are resorted to as a means of decomposition. I employ it in my guicantignition apparatus. (335.) At high temperatures, it is acted upon by the alkaline hydrates, and by almost all metals, especially tin and lead. I had a platinum crucible perforated, by fusing in it some flint glass, which can sists mainly of lead, silicic acid, and potash.

1593. The equivalent of platinum is 99.

Of the Compounds of Platinum with Ozygen.

of oxygen, which may be obtained from the chloride, by the addition of potash it forms also a bioxide, containing two atoms of oxygen to one of metal, as the same implies. The protoxide acts as an oxybase only, the bioxide, both as an oxybase and oxacid. In the last mentioned capacity it enters into combination with animalia, in the compound called fulminating platinum, and which we may with property call platinate of ammonia, or ammonium. Dr. Thomson alleges the existence of some other oxides of platinum.

Of the Compounds of Platinum with the Halogen Class.

regia above described, being in the state of a bichloride and acting as an acid agreeably to my fundamental definition, (631.) is capable of combining with other chlorides acting as chlorobases. With either the chloride of potassium, or the chloride of ammonium, (sal ammoniac, 1109.) it forms compounds which are but very sparingly soluble in water. Hence the precipitate resulting from the addition of the last mentioned chlorobase, and the employment of the chloride (or chlorobase) of potassium, in the precipitate as an acid, it is proper to designate it as a chloroplatine acid. In this I am supported by the authority of Dr. Thomson. It follows that the precipitates obtained as above described, are severally chloroplatinates of ammonium, and potassium.

1596. The superior solubility of the chloroplatmate of sodium, cushes us to distinguish solutions in which this metal exists as the radical, from those in which potassium performs the same part; as with the latter only is orange-coloured precipitate obtained, on adding chloroplatmic acid.

1597. Of Chloroplatinous Acid.—This name is given to the protochloride of platinum, as it is, according to Berzelius, capable of combining with the same chlorobases as chloroplatinic acid. Chloroplatinous acid is obtained by exposing the bichloride (chloroplatinic acid) to heat. It is alleged to have a grayish colour, and to be insoluble in water. Its composite with chlorobases must consistently be called chloroplatinites.

Experimental Illustrations.

in the malleable state. Precipitated from its solution bloride of ammonium, and chloride of tin. A precipiproduced in salts of potash by chloride of platinum, and salts of soda. Combustion of plan with tinfoil.

e Nomenclature of Compounds formed with Halogen Bodies, called Double Salts by Berzelius.

39. In order to present an intelligible view of the discordant names of alts above described, I will here subjoin a table of the names of some ounds formed with chlorine by platina, of which mention has been . (1596.)

of the various Names given to the Double Chlorides, such as those described in the case of Platina.

Names according to the old Theory of the Muriates.

•	•		•		
oniacal, or Ammonia,	}	Mu	riate of		Platinum.
Nan	res accor	ding to	Brande	•	•
					C Potaggium.
o bichloride of	•	-	-	•	Potassium. Sodium. Ammonium.
	Rerzeli	ian Nar	nes.		
	201 201				4 Ammonique.
ure platinico,	•	•	•	•	Ammonique. Sodique. Ammonique.
es according to Thom adoption					aware of their
					(Potassium.
platinate of	-	-	•	•	Potassium. Sodium. Ammonium.
lompounds formed was Berzelius designation	ith the I	Protochl follows:	oride,		
	•				(Potassique.
ure platinoso,	•	•	•	-	Potassique. Sodique. Ammonique.
re in this Compendiu	m design	nated h	u me .—		
s as sees compension	m ucoeg	milu o	<i>y m</i> e,—		(Potassium.
oplatinite of	•	•	-	•	Sodium. Ammonium.

1600. In order to have the nomenclature of the analogous compounds, in which other chlorobases or chloracids are introduced, it is only requisite

to change the corresponding epithet in the formula.

1601. By changing the syllables indicating the halogen ingredient, the nomenclatures of any of the double salts formed by any of the halogen bodies, may be seen, agreeably to the language of Brande, of Berzelius, or of Thomson and myself.

Of Bromides, Iodides, and Cyanides.

1602. Bromine forms with platinum a compound analogous in composition to the bichloride of that metal, and which, from its chemical properties, is entitled to the appellation of bromoplatinic acid.

1603. Iodine and fluorine both form compounds with platinum. The fluoride of platinum acts as an acid; the properties of the iodide in this respect are unknown.

1604. Cyanogen forms two compounds with platinum. The percyanide combines with cyanobases as an acid, and of course may be designated as cyunoplatinic scid.

Of the Compounds of Platinum with Sulphur.

1605. Platinum combines with sulphur in two proportions. Both sulphides combine as sulphobases with sulphacids, but the persulphide unites as a sulphacid with sulphobases of the alkalifiable metals.

Of the Power of Platinum, and other Metals in a divided or spongy form, to induce Chemical Reaction.

1606. In the spongy form in which platinum remains after the chlorine and ammonia of the chloroplatinate are expelled, it has the wonderful power of causing the inflammation of a mixture of hydrogen and oxygen gas. I have ascertained that this power is acquired by asbestos, porcelain earth, and charcoal, merely by scaking them in a solution of platinum, in aqua regia, and subsequent desiccation and ignition. Thenard states, that platinum filings, platinum leaf, or an association of fine platinum wires, exercise, in a greater or less degree, the same power as platinum sponge.

1607. The pulverulent mass, obtained by precipitating platinum by zinc, becomes incandescent in the vapour of alcohol. As the best means of obtaining platinum is that state of minute division in which it is most efficacious in producing this result, Liebig recommends that the chloride should be dissolved in a lixivium of caustic potash with heat; and that while the resulting liquid is still hot, alcohol should be added in small quantities, stirring the mixture until an effervescence arises from the extrication of carbonic acid. This, however, becomes so active as to render a very capacious vessel necessary for the process. The platinum precipitates in the form of a black powder, which is to be separated from the liquid, and washed successively with alcohol, with a solution of potash, with chlorohydric acid, and four or five times with water. When dried, the powder resembles lampblack, and soils the ingent Nevertheless, it consists only of platinum in a state of minute division, since it may be heated to a cherry-red in the air or in oxygen, without losing weight or undergoing any change in its properties. These are, however, destroyed by incandescence. which restores its metallic appearance. Under the burnisher, it becomes slightly of a metallic gray. In aqua regia, it dissolves without leaving any residue.

with a development of heat; and if, after being deprived of air and moisture by exposure to a vacuum over sulphuric acid, the atmosphere be rapidly admitted, it becomes red-hot. It causes the combustion of hydrogen or alcoholic vapour, when a contact with them with access of air, and becomes incandescent on falling on a surface wet with alcohol. If moistened with alcohol, it converts it, at the expense of the oxygen of the air, into acetic acid and acetic ether. Platinum, however prepared, gradually loses the property of causing a union between oxygen and hydrogen; but this property disappears much more rapidly when exposed to the action of the air, than when protected from its influence. Spongy platinum, moreover, remains effective longer than platinum under any other state; unless that in which it is procured by precipitation by zinc be an exception. Platinum leaf, which in the air is rendered powerless in a few minutes, when in the form of a scroll and accluded in a close vessel, remains effective for twenty-four hours. The power of producing a union between hydrogen and oxygen, may be restored by immersion as

an acid, or by incandescence

silver. 297

1609. Strips of sheet platinum, after being well cleansed by exposure as the electrodes of a Voltaic series; or by exposure, with certain precautions to acids, were

found by Faraday to cause the union of the elements of water.

1610. Platinum is not the only substance which possesses the property of protucing the combination of oxygen and hydrogen. Gold, precipitated from its solution by means of zinc and subsequently heated to redness, if assisted by a temperature of 122°, causes the union of hydrogen and oxygen. Osmium slowly produces the same effect, at a heat a little below that which is necessary in the case of gold. Spongy nickel acts slowly at the ordinary temperature of the atmosphere. Palladium, thodium, and iridium produce the same effect on a mixture of hydrogen and oxygen as platinum, though with less intensity. Charcoal, porcelain, glass, and rock crystal produce a union between hydrogen and oxygen, at temperatures lower than that at which it would otherwise take place.

Experimental Illustrations.

1611. A mixture of hydrogen and oxygen inflamed by platinum sponge, or platinated asbestos. Incandescence of platinum powder produced by moistening it with alcohol.

SECTION III.

OF SILVER.

1612. Silver exists in nature nearly pure, but usually containing a minute quantity of gold, copper, arsenic, or iron. It also exists in alloys, containing various equivalent proportions of arsenic, antimony, tellurium, or gold. It is found, likewise, in the state of chloride, iodide, sulphide,* and carbonate, and in a variety of galena, the native sulphide of lead, called, in consequence, argentiferous galena. In consequence of its fusibility and insusceptibility of oxidizement,—when any metallic alloy, containing t, is exposed to intense ignition with access of air, the silver is separated rom any metal which, when thus exposed, is liable to be converted into an oxide. In the small way, this object is effected upon the cupel, in the operation called cupellation. A cupel is a small flat cylinder, made of bone arth obtained by calcination, in the upper surface of which there is a henispherical cavity. In this cavity any gold or silver to be refined is placed vith a portion of lead. The cupel is then placed in a small earthenware wen, called a muffle, and exposed to a heat sufficient to render and keep he metals fluid. Under these circumstances, the lead is oxidized and vitriied, and promotes a similar change in any other metals present, which are usceptible of oxidizement under the circumstances in question. The vitrems matter, thus produced, is absorbed by the bone earth. After the process nas endured sufficiently, only the gold and silver, or other noble metals, should any be present, can remain upon the cupel. This stage of the prozess is indicated by the metallic surface being no longer obscured by any ilm of oxidized matter. As in the cases in which this process is employed, 10 other noble metals are liable to be present besides gold and silver, I shall reat of it only in reference to them.

[•] The student is requested to recollect that sulphide and sulphuret are synony.

1613. From the alloy purified upon the cupel, the silver may be sensored by nitric acid, when the gold does not exceed a fourth of the whole. Is order, therefore, to enable the nitric acid to dissolve the silver, the man is fused with the addition of as much of this metal, as will establish the requisite ratio between it and the gold. This is called quartation. The process of separating the metals afterwards by nitric acid, is called partiage

up. The silver will precipitate as a chloride, and the parting will be effected the easier, in proportion as the quantity present of the last mentioned metal

is leas.

of lead, in the process called liquation. The alloy is fused with two and a half parts of the metal last mentioned, and east into thick round cakes. These are subsequently exposed in a reverberating furnace, to a heat maticient to melt the silver and lead, leaving the copper, which has warrif any affinity for lead. The silver is afterwards separated from the lead by

cupellation.

The white crystals spontaneously afforded by a solution of the coin in number acid, cautiously drained, and washed with a portion of water barely subcient to remove every vestige of green, yield a solution of pure silver. The residual liquid may be decomposed by copper, and the precipitate reducibled, and crystallized by evaporation; and thus a fresh crop of whate crystals may be procured. The whiteness of the crystals may be decomposed the criterion of their purity. Silver precipitated by mercury, as in the case

of the arbor Dianæ, only requires ignition to render it pure-

degree of metallic brilliancy; and next to gold, it is the most mailcable and ductile metal. In metallic whiteness, it is pre-eminently beautiful, and at tenacity inferior only to iron, copper, and platinum. Its specific gravity is 10.5, and equivalent 108. It is the best conductor of caloric, fixes at a low white heat, is as difficult to oxidize in the fire as gold, but is more lather to tarnish when exposed to the atmosphere, from its susceptibility to the action of sulphur and chlorine. Hence it is blackened by eggs and by salt water.

1618. By the compound blowpipe, electricity, or galvanism, silver a

fused, oxidized and dissipated.

1619. Exposed to nitric acid, it is oxidized by one portion, and district by the other. In fact this acid is its proper solvent. The resulting mate, when fused and cast into sticks, forms the lunar caustic of the shops. It consists of one atom of acid 54, and one of oxide 116, = 170.

1620. Sulphuric acid, when cold, has no reaction with silver. At a boiling heat, the metal is oxidized at the expense of one portion of the cold and the oxide, thus formed, is dissolved by another portion, as in the cold.

of nitric acid.

1621. Silver combines with phosphorus, and in minute proportion with carbon and silicon.

Of the Compounds of Silver with Ozygen

1622. Silver forms two oxides. The protoxide is obtained by decomposing the nitrate by potash or soda. It is of a deep olive colour, stability soluble in wait, and, according to Thenard, sufficiently alkaline to render syrup of violets grown. It revives simply by the influence of heat, and of course is reducible when heated with those radicals, which, under the same circumstances, combine, per se, with exygen-

SILVER. 299

When thus reduced, it must of course produce the vivid ignition which is consequent

to the presence of pure oxygen.

1623. This oxide, by uniting with ammonia, produces a fulminating compound, so dangerous that few persons have been willing to encounter the risk of making it. This should not be confounded with the fulminate of silver, consisting of the same

metal and fulminic acid. (1312, &c.)

1624. The peroxide of silver is formed around the wire, proceeding from the positive pole of the Voltaic series, when a weak solution of nitrate of silver is placed in the circuit. It crystallizes in long needles, endowed with a metallic brilliancy. It does not combine with acids, but when presented to them, by a partial relinquishment of oxygen, passes to the state of protoxide. This oxide does not act either as a base or an acid. It detonates with phosphorus, if struck with a hammer while in contact with that substance; and when subjected to ammonia, disengages nitrogen from it by oxidizing the hydrogen.

1625. The protoxide of silver consists of one atom of silver, equivalent 108, and

one of oxygen, equivalent 8 = 116.

1626. Some chemists suppose the existence of a suboxide of silver.

Of the Compounds of Silver with the Halogen Class.

1627. Silver unites with chlorine when heated in it, or presented to it in solution. The resulting chloride is one of the most insoluble combinations. Hence, silver is not soluble in aqua regia, or any other liquid containing chlorine; and on this account, soluble chlorides yield a precipitate, when solutions of silver are added to them.

1628. The chloride of silver is white and tasteless, and, according to Thenard, when exposed to light, is decomposed, forming a subchloride. It combines as a base with the electro-negative, and as an acid with the electro-positive chlorides.

1629. It is susceptible of fusion without decomposition, forming what was called

by the old chemists, luna cornea, or horn silver.

1630. Chloride of silver is soluble in liquid ammonia, forming chloroargentate of ammonium. It may be decomposed by hydrogen, if brought into contact with this gas in its nascent state, as when evolved from zinc or iron by a diluted acid. It is easily decomposed by the compound blowpipe, supplied with hydrogen and atmospheric air; also by fusion with the fixed alkalies, or when boiled in water with shreds of iron. The formation and subsequent decomposition of the chloride, is one of the modes of obtaining pure silver.

1631. The chloride of silver is composed of one atom of silver, and one atom of

chlorine.

1632. Bromine and iodine form, with silver, compounds analogous in composition to the chloride. The bromide and iodide, acting as acids, combine with the bromides and iodides of the alkalifiable metals, acting as bases. A native iodide of silver has been found in Mexico.

1633. Fluorine and cyanogen both combine with silver; the fluoride acts as a

base, the cyanide as an acid-

Of the Compound of Silver with Sulphur.

1634. The sulphide of silver is solid, ductile, easy to cut, of a lead-gray colour, with a metallic brilliancy, more fusible than silver, crystallizable in cubes or octo-

hedra, and indecomposable by fire. It acts as a powerful sulphobase.

1635. This sulphide is produced whenever silver is exposed to sulphur, or sulphydric acid. The blackening of silver spoons by eggs, is ascribed to the existence of a minute portion of sulphur in the albumen. The impression has been entertained that persons, who use the water of the white sulphur springs freely, find silver coin, carried about them, blackened by the sulphur introduced into the blood. This, however, may arise from their frequenting the spring, and thus exposing the silver about their persons to the action of the sulphydric acid gas which is continually evolved from the water. I am, however, far from deeming the fact improbable. It would, a priori, be no more surprising that the animal frame should be imbued with sulphur than with mercury.

Experimental Illustrations.

1636. Exhibition of an assay furnace and muffle; also of a cupel. Oxidizement and solution of silver in nitric

acid exemplified; also its precipitation by chlorides, phosphates, arsenites, arseniates, copper, and mercury. Fused subnitrate of silver, or lunar caustic, exhibited. (1619.)

SECTION IV.

OF MERCURY.

1637. This well known metal is found in nature in the metallic state, pure, and in union with silver; also in the states of sulphide and of chloride. It is obtained principally from the native sulphide, cinnabar, the most abundant and prolific of its ores, by distillation with the hydrate of lime. In this country it may always be procured nearly pure, in the iron bottles in which it is imported. It is alleged that mercury may be purified by distillations from iron filings. I once distilled several hundred pounds in this way for my mercurial reservoir, but did not find it to be quite pure afterwards. A better mode is to digest it at a heat below the boiling point, with dilute sulphuric acid in a glass, porcelain, or stoneware vessel. Under these circumstances, any metal having a greater affinity for oxygen than mercury, will be taken up by the acid.

1638. If on being agitated in a cup of white porcelain mercury does not soil the surface, it may be considered as

pure.

1639. When triturated or violently shaken with other matter, so that this may be sufficiently interposed between its particles to prevent them from touching, it is liable to be comminuted into a black or deep blue powder. According to Berzelius, it is in this form that mercury exists in blue ointment. It follows that the method which has been recommended for the purification of this metal by agitating it so as to cause the oxidizement of the more electro-postive metal which it may contain, is pregnant with the evil that a portion of the metal separates with the resulting oxides. There is a large proportion of mercury separates with the lime, during the oxidizement of the radical of this earth in the amalgam of calcium.

1640. Properties.—Mercury is the only metal which is liquid at the ordinary temperature of the atmosphere.

and brilliancy it resembles and rivals silver. It into a malleable solid at —39°, and boils at 665°. some experiments by Faraday, it would appear that ry vaporizes to a minute extent, whenever the temperature of the air is above 70°. At the temperature of specific gravity is 13.6; but in freezing, it is ind to 14.4.

1. Mercurial compounds are all volatilizable by heat; ercurial salts, when moistened and rubbed upon cop-

ver it with a film of mercury.

2. It is alleged that if a drop of any liquid containercury be placed upon gold, and touched with the of a knife, or a piece of iron wire, the mercury will cipitated upon the gold.

f the Alloys of Mercury with other Metals, called Amalgams.

3. All metals combine with mercury, directly or iny. Its compounds have the generic name of amal-

In the case of gold, silver, zinc, lead, tin, and bisthe amalgamation is rapidly effected. It is less produced with copper, unless when this metal sepanercury from the acids. It is difficult to unite mervith platinum, and still more so with iron, owing, ly, to the great difference in fusibility.

1. Mercury unites energetically with the metals of calies. In the case of sodium, a species of combususues, so that the mass becomes red-hot; and when
usues is thrown upon mercury, it is repelled with vio-

and with a disengagement of light.

5. The amalgam of potassium is an efficient instruin the evolution of the amalgam of ammonium.

5. The equivalents of mercury, and of its compounds xygen, chlorine, and sulphur, are as follows:—

ry,	-	•	•		equivalent	202
ry, ide,	1 atom	mercury	with	1	atom oxygen	210
e,	1	"			atoms "	218
hloride,	1	"	"	1	atom chlorine	238
ride,	1)			atoms "	274
mlphide,	1	77	77	1	atom sulphur,	218
hide,	1	??	77	2	atoms ,,	234

Of the Compounds of Mercury with Oxygen.

1647. There is still some discordancy in the opinions and language of chemists respecting the oxides of mercury. It has been generally held that there are two oxides of this metal; one black, the other red. According to the table of equivalents in the Appendix, the atom of mercury = 202, with one atom of oxygen, forms the black oxide or protoxide, = 210, and with two atoms of oxygen forms the red oxide, peroxide, deutoxide, or bioxide, = 218. Dr. Thomson, conceiving the atom of mercury to have only half the weight here assigned to it, considers the black oxide as consisting of two atoms of metal, and one of orygen, and consequently designates it as a suboxide. The difference in this case, therefore, is only hypothetical. But Guibourt has advanced that there is only one oxide of mercury, the black oxide being composed, according to him, of the red oxide and metallic mercury. Donovan, or the other hand, has shown that, when a small quantity of calomel is added to a comparatively large quantity of potash, a pure protoxide or suboxide is obtained.

1648. The protoxide of mercury may be obtained by digesting the protochloride with an excess of potash. The oxygen and chlorine exchange places, and the chloride of potassium, and protoxide of mercury are formed. This oxide may be precipitated from a solution of the nitrate of the protoxide, by the addition of an alkaline solution. It has likewise been supposed to be produced when mercury is subjected to long continued agitation, in contact with air. It is, however, alleged by Berzelius, that the black powder, thus obtained, is metallic mercury in a state of extreme division, to which it could not be reduced without the interposition of the oxide, resulting from the presence of a more oxidizable metal; that, when the metal is quite pure, and free from the interposition of heterogeneous particles, it undergoes no change by agitation; but that if, under the same circumstances, it be triturated with grease, gum, powdered glass, or sand, it may be reduced to a black metallic powder. He conceives that all the preparations made by triturating mercury with greass, gum, or other viscid substances, contain the metal in a divided state, but not oxidized.

1649. The protoxide of mercury is a black powder, which

y exposure to light, or to the heat of boiling water, is onverted into metallic mercury and bioxide.

1650. The bioxide may be procured by long exposure of nercury to a heat sufficient to cause a gentle ebullition, he air having free access. It may likewise be obtained y expelling the acid from the nitrate by heat. Berzelius norms us, that, agreeably to the opinion of some respect-ble physicians, it is only when procured by the former nethod that it is fit to be used internally. He attributes his difference to the fact of its sometimes retaining, when btained from the nitrate, a small portion of nitric acid. This oxide, when in mass, is of a beautiful red colour, but, when powdered, assumes a yellowish appearance. It is ecomposed into metallic mercury and oxygen at a heat little below redness. In water it is slightly soluble, and is solution has the property of rendering syrup of violets reen.

1651. Both the oxides of mercury act as bases. The ioxide forms with ammonia a fulminating compound.

Of the Reaction of Acids with Mercury and its Oxides.

1652. When nitric acid, whether cold or hot, concentrated or dilute, is brought into contact with mercury, one portion of the acid is decomposed, imparting oxygen to the netal; the oxide thus formed being dissolved by the renainder of the acid. When the metal is in excess, the rotoxide is principally formed. When the acid is in excess, the bioxide predominates. Usually, more or less of ach oxide is formed.

1653. Whether concentrated or dilute, cold sulphuric cid does not react with mercury; but when the concentrated acid is boiled on the metal, the phenomena are analogous to those which ensue in the case of nitric acid. One ortion of the acid yields oxygen to the metal; another ombines with the oxide thus created.

1654. As in the case of nitric acid, we may have the xybase of mercury in the state of bioxide, or of protoxide, accordingly as the acid or the metal is in excess, or as the time allowed for oxidizement is greater or less.

1655. Each oxide of mercury forms three salts with nitric acid. When washed with hot water, the bisulphate of the bioxide yields a yellow compound known under the name of turpeth mineral, which, according to a recent ana-

lysis by Kane, is a subsulphate, consisting of one atom of neutral sulphate, and two atoms of protoxide, the formula

being HGO SO³ + HGO.

nerally no reaction with acids; yet in the state of an oxide, it is no doubt liable to be combined with any of them. About one hundred mercurial salts are mentioned in Thomson's Inorganic Chemistry. With these compounds, without a special motive, it would be worse than useless for a medical student to burthen his memory. But it is fortunate that, in this branch of our knowledge, we are aided by analogy, and that we are enabled, when we hear of an oxacid, to infer that the formation of a corresponding salt with each oxide of mercury is possible. Moreover, agreeably to the received principles of chemical nomenclature, we are enabled to assign to the compound thus imagined, a name which would be recognised by another chemist.

1657. In the metallic state, mercury has no reaction with acids, having hydrogen for their radical, called hydracids, by some chemists; but which are, in this book, designated as halohydric, or amphydric acids. (660.) Yet, excepting in the case of the earths proper, it may be assumed that, when any metallic oxide is presented to any of the hydracids, water will be formed by the oxygen of the one, and the hydrogen of the other; while the metal and basacigen body will unite, and, in a majority of instances, the metal will acquire the same number of atoms of the basacigen body, as it relinquishes of oxygen. of course, that on subjecting a mercurial oxide to chlorohydric, bromohydric, iodohydric, fluohydric, cyanhydrc, sulphydric, selenhydric, or telluhydric acid, a chloride, bromide, iodide, fluoride, cyanide, sulphide, selenide, or telluride, will generally, if not universally, result.

Of the Chlorides of Mercury.

1658. When a solution of chloride of sodium and of nitrate of the protoxide of mercury are mingled, the oxygen of the oxide, and chlorine of the chloride exchange places; so that protochloride of mercury (calomel) precipitates, while the oxygen and sodium uniting, remain in solution, and in union with the nitric acid.

1659. If the solution of chloride of sodium be added to

on of the nitrate of the bioxide, the two atoms of in this oxide will exchange places with the chlotwo atoms of the chloride; so that the mercury will, atoms of oxygen, acquire a like number of atoms rine, and be thus converted into one atom of bie, or corrosive sublimate, which will remain in soluthere be enough water present. Thus the quantity rine transferred is regulated by the quantity of oxythe oxide employed; the protoxide producing the doride, the bioxide the bichloride. (1657.)

· The complex affinity which causes these changes, s either in the wet or dry way; that is, whether stances be mixed in solution, or sublimed together. sulphate of the bioxide of mercury produces these when sublimed with certain compounds of chlochloride of sodium for instance. Corrosive subs thus procured; and, by trituration with mercury, id sublimation, and washing in boiling water, may erted into calomel; or the bisulphate, by trituration further portion of the metal, being converted into alphate, forms calomel directly by sublimation with n salt.

. The process for the manufacture of the proto-3, has been improved by causing its nascent vapour ningled with steam, which, interposing between the s, prevents them from adhering as they condense. ire thus obtained in a state of more minute divian could be effected by trituration, and the aqueous s, in condensing, combine with and remove any s of corrosive sublimate which may be generated neously with those of the calomel. Calomel thus d, has been distinguished as Howard's hydrosub-

Le Chlorine does not combine with mercury in the innode abovementioned only. A jet of chlorine burns

eously in mercurial vapour, forming a bichloride.

The chlorides of mercury may likewise be obby subjecting the oxides to chlorohydric acid; in case the hydrogen of the acid, and oxygen of the orm water, while the mercury and chlorine unite; toxide giving rise to the protochloride, the bioxide bichloride, as already explained. (1657.)

. The processes for manufacturing these important

compounds of mercury, are very numerous. They have, however, but one object—that of presenting chlorine and mercury to each other in due quantities, and intimately mingled. When the chlorine is in excess, corrosive submate is formed; when the metal predominates, calomel.

usually more compact than corrosive sublimate. It is tasteless, inodorous, and unalterable by exposure to the atmosphere if protected from light; but by this it is blackened, and partially reduced to the metallic state. It is blackened by alkaline solutions, by the generation of protoxide; and when the surface is removed, it appears yellow, so that a scratch made with the nail, is productive of a yellow streak. It is less volatile than the bichloride. This chloride acts as a base.

1666. The bichloride or corrosive sublimate is white, more or less crystalline, and transparent. It is soluble about twenty parts of cold water, but more so in be water, whence crystals are obtained by refrigeration. dissolves in two parts of alcohol, and in three parts of ether, by weight. It is not soluble either in sulphuric or nitric acid. On the application of heat, it sublimes changed. When added to an alkaline solution in excess, a yellow hydrated bioxide of mercury precipitates. The proportions being reversed, as when an alkaline solution is dropped into the solution of this chloride, a compound of the bioxide and bichloride is precipitated, which is of a brick-red colour. Berzelius designates this compound as the chlorure mercurique basique, while Thenard gives it the name of bioxido-chlorure. The latter appellation, changed to oxychloride,* in order to render it consistent with the nomenclature adopted in this work, and more easy to pronounce, I shall employ for this, and for analogous compounds formed with other metals.†

1667. Ammonia throws down from a solution of the bichloride, a compound called ammoniated mercury in the

^{*} I believe this name is now generally preferred.

[†] This compound probably furnishes one among many other instances, in which an electro-positive compound of one basacigen body, unites with an electro-negative

compound of another.

It might indeed be alleged, that in this case the metal acts the part of a bearing body, agreeably to my definition, (623,) as it enters into two compounds, one electronegative, the other electro-positive, which form a tertium quid. This view of the subject corroborates the remark, which I have elsewhere made, that nature has all fitted her bodies for distinct classification; and that consequently there will be coming in which some of the bodies associated in one class, will appear to belong to another.

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ted States' Pharmacopæia, but more generally known hydrargyrum precipitatum album, or white precipitate. ne light, derived from organic chemistry, has lately led a new view of this preparation, which it will be more venient to present, in treating of that branch of the nce.

- 668. The bichloride of mercury has a most nauseous allic taste, and is a virulent poison. The best antidote it is albumen, which may be given in the form of the te of eggs, diluted with water. This chloride, acting an acid, combines with the chlorides of ammonium, assium, sodium, barium, and magnesium. As a base it abines with chlorohydric acid, forming a crystalline apound, which effloresces and is decomposed when exed to the air.
- 669. The salt, long known under the antiquated appelon of sal alembroth, and formed by mixing solutions of ammoniac and corrosive sublimate, is now by Berzelius sidered as a double salt, and by Bonsdorf, a chlorohygyrate. This last mentioned name accords with the nition of acidity and basidity which I have proposed.

Of the Bromides, Iodides, Fluorides, and Cyanides of Mercury.

370. As respects the means of their production, the ratio of their conent atoms, and their qualities in general, there is the greatest analogy cen the compounds formed by mercury with the halogen class. Hence, ag treated particularly of the chlorides, I shall treat with the utmost ty of the compounds named at the head of this article.

571. Bromine forms two compounds with mercury. The protobromide hite, pulverulent, and insoluble. It is obtained by the reaction of the

nide of potassium with the nitrate of the protoxide of mercury.

372. The bibromide is formed by subjecting mercury to the action of nine and water. It is soluble, crystallizable, fusible, capable of volatilion, and colourless. With the alkaline bromides it acts as an acid.

- 173. The protiodide of mercury is obtained by the reaction of the le of potassium with the nitrate of the protoxide of mercury. It is olive ared, insoluble in water, and corresponds in composition with the prot-
- solution of the bichloride of mercury. A reciprocal decomposition place, and the bichloride of mercury and the chloride of potassium are red. The biiodide is fusible, volatile, and of a transcendently beautiful let colour. As an acid it combines with the iodides of many of the he and alkalies, and with the iodides of zinc and iron. As a base it with iodohydric acid. It consists of one atom of mercury, comd with two atoms of iodine.

1675. Another iodide exists, containing less iodine than the biiodide, but

more than the protiodide.

1676. Although cyanogen does not combine with mercury directly, a bicyanide is obtained when the bioxide of this metal is digested in water with Prussian blue. I shall more fully explain this process when treating of iron.

1677. The employment of the bicyanide in the evolution of cyanogen and cyanhydric acid, has already been mentioned. (1296). The bicyanide of mercury forms combinations with the alkaline cyanides, in which it plays

the part of an acid.

1678. The fluoride of mercury is obtained by the action of fluohydric acid on the bioxide. It acts both as an acid and base. It is yellow, volatile, and when volatilized in platinum or glass vessels, corrodes them. If subjected to water, it is resolved into a soluble and an insoluble compound.

Of the Compounds of Mercury with Sulphur.

1679. Agreeably to the list of equivalents near the beginning of this section, mercury forms with sulphur, a protosulphide and a bisulphide. When a weak solution of the nitrate of the protoxide of mercury is impregnated with sulphydric acid, the protosulphide is produced in the form of a black precipitate. It appears to be an unimportant compound. The protosulphide acts as a sulphobase.*

1680. The bisulphide may be generated by impregnating with the same gas a solution of the bichloride. Thus obtained, it resembles the protosulphide in assuming a black colour, which, however, may be changed to red

by sublimation.

1681. The bisulphide (artificial cinnabar) is procured in the large way. by fusing one part of sulphur, stirring in gradually six or seven of mercury. and subjecting the resulting black mass to the process of sublimation in close vessels.

1682. Kirchoff procured cinnabar by a long continued trituration of mercury and sulphur with a solution of caustic potash, aided by a gentle

heat.

1683. Bisulphide of mercury is attacked neither by sulphuric, nitre nor chlorohydric acid, nor by caustic alkaline solutions; but when subjected to chlorine, either in aqueous solution as in aqua regia, or in the gaseous form, it is converted into bichloride of mercury, and bichloride of sulphur.

1684. Berzelius alleges that this bisulphide performs the part of a base in combining with the aëriform sulphacids. It also forms compounds with

the bichloride, bibromide, biiodide, and bifluoride of mercury.

1685. Equal parts by weight of mercury and sulphur, triturated together, form a black mass, called, from its colour, *Ethiops mineral*, which is now considered as a mixture of bisulphide of mercury and sulphur.

[&]quot;Guibourt, who has been mentioned as questioning the existence of more than one oxide of mercury, has alleged his disbelief in the existence of more than one mercurial sulphide; the black sulphide being, in his opinion, a mixture of the set with the metal. Seftstrom controverts this allegation, and Thenard, citing be opinions and that of Guibourt, inclines in favour of those of Seftstrom. Thenard alleges that at least the protosulphide exists as a base, in combination with sulphides. acting of course as sulphacids agreeably to my definition. (631)

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Of the Phosphurets of Mercury.

1686. In its habitudes with mercury, phosphorus displays that analogy with sulphur which is in general so remarkable. It forms, according to Berzelius, a red and a black phosphuret. The latter results from the digestion of bioxide of mercury with phosphorus and water; the former, from exposing phosphorus to the vapours of the bichloride of mercury.

Experimental Illustrations.

1687. Ebullition and distillation of mercury. Its compounds with oxygen and sulphur, exhibited. Action of nitric acid and of sulphuric acid on the metal. salts, subjected to hot water. Black oxide and red oxide, severally dissolved in nitric acid. Chlorohydric acid precipitates calomel from the one, but occasions no precipitate in the other. Fixed alkalies and alkaline earths produce a black precipitate in the nitrate of the black oxide or protoxide, an orange precipitate in the nitrate of the red oxide or bioxide. Similar results obtained by adding them to calomel and corrosive sublimate; the former giving the black, the latter the red oxide. Hydrargyrum precipitatum album, precipitated from solution of bichloride by ammonia. Solutions of the different mercurial oxides precipitated by iodide of potassium. Inflammation of chlorine with mercurial vapour. Explosion of fulminating mercury. Diversity of precipitates produced by adding bichloride to an excess of alkali, or adding the latter to an excess of bichloride. (1666.)

Combustion of Mercury with Chlorine.

1688. This experiment may be performed by means of the apparatus represented by the following cut. Let there be a glass globe, furnished with a neck and tubulure, and holding about two gallons of chlorine. Into the neck, let a trumpet-shaped tube, B, reaching to the bottom, be fastened air-tight by means of a cork.

16:9. Let another tube, about fifteen inches in length, and tapering towards one end, so as to form a capillary orifice, be fastened, at the other end, into the lateral tubulure of the globe. Provide a globular receiver, R, with a neck on one side, and

a perforation on the other, opposite the neck.

1690. Let the lower part of this vessel be occupied by about a gill of mercury, and exposed to a chauffer of coals, so as to fill the whole cavity of the vessel with the vaporized metal. Under these circumstances, introduce the pipe, D, proceeding from the lateral tubulure, into the neck of the receiver, so that the capillary orifice may be near the perforation; and immediately afterwards pour chlorohydric acid into the tube, B. This will subject the chlorine to pressure without absorbing it, and consequently cause it to escape in a jet from the capillary orifice in the pipe. Hence, mingling with the vaporized mercury, it will produce a feebly luminous flame.

1691. Instead of using the globe and its appendages, I have, in a majority of instances, employed a tubulated retort, with a long narrow beak, for the production of the jet of chlorine. The retort being sufficiently supplied with manganese, and a



glass funnel with a cock being fastened into the tubulare, as much rhioroby acid is allowed to enter through the funnel, as will generate a sufficient quantity objection to produce a jet from the capillary ornice, in which the beak of the result is purposely made to terminate. The beak of the retort being made to occupy the place of the tube, represented in the figure as proceeding from the globe, the expriment is, in other respects, the same as that above described.

1692. Since this engraving was made, for the purpose of supplying chlorum, I have found my self-regulating reservoir of chlorine, to render the perfermence of this experiment more convenient and less precarious than the apparatus alors re-

presented. (788, 975.)

SECTION V.

OF COPPER.

1693. Copper is occasionally found in nature in the metallic state; also in those of oxide, carbonate and sulphide. It is obtained principally from the sulphide. The sulphide being acidified and volatilized, and the metal oxidized by torrefaction, the resulting oxide is decomposed by heat and charcoal.

1694. The copper of commerce contains, according to Berzelius, a minute portion of sulphur and carbon. It may be purified by solution in concentrated and boding chlorohydric acid, and subsequent precipitation by a bright plate of iron.

1695. Properties.—The lustre and peculiar colour of this metal are too well known to need description. Es-

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titanium, it is the only red metal. It is very maland ductile, and next to iron in tenacity. It fuses hite heat. Alloyed with a small quantity of tin, forms bronze; with a larger quantity, bell metal. with zinc, or subjected to the vapour of this metal, ved from calamine when heated with charcoal, it is ted into brass.

Heated to bright redness in a tube, the gas is desed, and the copper becomes brittle, although its is but slightly, if at all increased. This change is ed to be due to a different aggregation of the partithe copper, induced by the formation and subse-

lecomposition of a nitruret of copper.

- '. The blade of a knife, or any bright piece of iron l, is a test for copper in solution; as a film of this vill be precipitated upon the iron or steel and comte to it the appearance of copper. Ammonia, added in excess, produces a blue colour in water ing a very minute quantity of copper, (523, &c.) ave ascertained that it requires twice as much copproduce a blue tinge with ammonia, as to produce the cyanoferrous acid of the cyanoferrite of potashe appropriate hue of the cyanoferrite of copper, s a peculiar rich reddish-brown. With this test I etected copper in the rain water, proceeding from ut of a copper roof.
- s, forming unimportant compounds. When present quantity, it has an effect upon this metal similar which carbon has on iron, rendering the copper tough for cutting instruments. Carbon and silicon mbine with copper.
- . The specific gravity of copper is nearly 9.
- The equivalents of copper, and of its compounds ygen, chlorine, and sulphur, are as follows:—

•	•	•	•	•	•	32
ioxide,	2 atoms co	pper with 1	atom ox	ygen,	•	72
protoxide,			atom	"	•	40
•		pper with 2	atoms o	xygen,	•	48
le,	2 atoms		atom ch		•	100
oride,	1 atom	,, 1	atom	"	•	68
ie,	2 atoms	,, 1	l atom su	lphur,	•	8 0
le, shide,	1 atom	,, 1	latom	"	•	48

Of the Compounds of Copper with Oxygen.

1701. There are three oxides of copper, a dioxide,

protoxide, and a peroxide.

dioxide; since it forms the dull red exterior coating a copper, as it comes to us from the manufacturer. When this oxide is subjected to liquid chlorohydric acid, a double decomposition ensues, and water and a dichloride of copperare formed, the latter remaining in solution. Subjected to nitric acid, the dioxide is converted into protoxide by comportion of the acid, and dissolved by the other. By it niting the protoxide with metallic copper, the anhydrous dioxide is produced. On mingling solutions of potash and of the dichloride, an orange-coloured hydrate of the distinct pound without partially converting it into protoxide.

colour. It is formed upon sheet copper, when exposed to a bright red-heat with access of air. To obtain it, however, Thenard recommends that the nitrate or sulphate be intensely ignited in a stoneware retort, by which means the acid is volatilized, and the protoxide remains. The oxide is formed when copper is dissolved in natric or appeared acid, and enters into combination with them, form

ing a nitrate, or sulphate.

1704. Thenard alleges that it is soluble in ammoniate only when combined with water or some acid; and that is insoluble in the fixed alkalies, whether hydrous or at

hydrous.

1705. That the fixed alkalies promote its oxidizence is evident; since sheet copper, or brass, moistened wit alkaline solutions, always becomes green in the air. think it probable, that carbonic acid co-operates in pro-

ducing this result.

the bioxide (deutoxide) of hydrogen with a weak solution of nitrate of copper, and adding just enough alkali to a compose all the nitrate. These conditions being realise a brownish-yellow gelatinous mass subsides, which, and being washed upon a filter with cold water, and dried the vacuum of an air pump over sulphuric acid, forms the peroxide. This oxide has no taste or smell.

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1707. The dioxide of copper acts feebly as a base, the rotoxide energetically, and the peroxide plays the part either of a base nor of an acid.

1708. Nitric acid diluted nearly to the specific gravity f 1.2, protoxidizes and dissolves copper, producing a blue plution, which yields, by evaporation, elegant blue crystals. The ignition which ensues when these crystals are alverized, moistened, and rolled up in tinfoil, has been aduced as an exemplification of the influence of water in romoting chemical reaction. (829.)

1709. The crystals of nitrate of copper are of a deeper lue than those of the sulphate, and are deliquescent. At moderate temperature, these crystals fuse, and lose a part

f their water of crystallization.

1710. Sulphuric acid, boiled on copper, oxidizes and displayes it without heat, as nitric acid does. The resulting ompound forms the blue crystals of the sulphate, called in ommerce blue vitriol or blue stone. (494.)

1711. A compound is obtained by triturating sulphate copper with carbonate of ammonia, called cuprum amoniatum (ammoniated copper) in the Pharmacopæias.

1712. This contains the ammoniacal sulphate of copper, rith some portion of the carbonate undecomposed. It may e a mixture of ammoniacal sulphate, and ammoniacal arbonate of copper. It is designated as ammoniated opper in the United States' Dispensatory; and the authors llege that there is some obscurity as to the mode in hich its ingredients are associated. It has been stated bove, that the protoxide of copper is not soluble in amionia, unless when united with water acting as hydric cid, or with some other substance capable of performing ne part of an acid. Berzelius mentions that the protoxle of copper may be kept in a bottle containing liquid mmonia, without tinging it blue; but that the introducon of only a few drops of any ammoniacal salt, the caronate for instance, causes the well known striking blue polour of the cupreous solution, formerly called aqua sap-He also alleges that cuprum ammoniatum conains the ingredients in such proportion, that the alkali aturates twice as much acid as the copper.

Of the Compounds of the Oxides of Copper with Acetic Acid.

1713. The oxides of copper form salts with almost every acid, whether mineral or vegetable. Among these, note are better known than its combinations with acetic acid, of which one is designated in commerce as verdigris, the other as crystals of Venus.

1714. Crude verdigris is a mixture of neutral acetate

and subacetate of copper, with some impurities.

1715. The neutral acetate crystallizes readily, and in the crystalline form has received the name of crystals of Venus; Venus having been one of the names given to copper by the old chemists.

1716. The subacetate consists of one atom of acid with two of protoxide, while the neutral acetate consists of one

atom of each constituent.

1717. The salts of the protoxide of copper are all of an intensely blue or green colour. This does not appear to be true in the case of the dioxide; since, according to Berzelius, when verdigris is subjected to heat, colourless crystals of the acetate of the dioxide sublime, so as nearly to fill the beak of the retort. Thenard alleges that when the hydrated dioxide is subjected to liquid ammonia, a colour-less solution of it results.

Of the Compounds of Copper with the Halogen Class.

prepared by the action of chlorohydric acid on a mixture of the protoxide of copper and finely divided metallic copper. From this chlorohydrate, the dichlorde may be precipitated by water. It is likewise produced by heating the protochlorice in conversels, by which one-half of the chlorine is expected. This chloride may be obtained from the protochloride, by digestion in chlorohydric acid with copper with or filings. When thus evolved, it subsides in crystals, which can only be end in vacuo over sulphuric acid, as in the air they are converted into a compounded the diexide with the protochloride, forming of contact an oxychloride.

1719 The dichlerale is soluble, crystallizable, and tusble by heat. It could

two atoms of copper, and one of chlorine.

with the protochaoride may be formed either by the reaction of chlorohydric with the protoxide, or by the combustion of copper wire or leaves in chlorine. It is crystallizable, fumble, of a bluish-green colour, and an astringent taste. It is crystallizable, fumble, and decomposable by heat. This chloride attracts in isture, and is very soluble in which therefore written with a solution of it, remain invisible intil freated, when the become yellow. It constitutes, of course, a species of sympothetic ink. The prochloride acts as an acid with the onlorides of potassizin and arimnemum, and with other chlorideses, forming with them can recapitates.

nard. These formed with reduce appear on the specific neighbor that With discussion forms two compounds, a protection and a perfluit rise. The products acts as a base, the perfluit rise both as a base and an acid. Cyanogen enters all the

bination with copper in two different proportions

LEAD. 315

Of the Compounds of Copper with Sulphur and Scienium.

A disulphide of copper may be produced by the fusion of the metal, or its its sulphur. It is found pure in nature, and likewise combined in definite us with the sulphides of antimony, arsenic, bismuth, and iron, in which sulphur is usually divided equally between the metals. Berzelius alleges tide to be a powerful sulphobase. Yet, in its combination with the sulphide t cannot be supposed to act as a base, as iron is more electro-positive than The affinity between the last mentioned sulphides is so energetic, that the sulphocuprate of iron cannot be decomposed by the united action of carbon ed alkali.

Pretosulphide of copper is formed when this metal is precipitated from its

by sulphydric acid.

Copper, by fusion with various sulphides of the alkalifiable metals, is made with several proportions of sulphur, but the resulting compounds are unim-

The union of copper with selenium is productive of heat and light; in which as in others, the analogy between selenium and sulphur is sustained. The compound is a diselenide. It is found in nature, but does not appear to interesting properties. A protoselenide is formed when copper is precipiselenhydric acid.

Experimental Illustrations.

6. Solution of copper in nitric acid, and its precipiby iron. Effects of ammonia; also of cyanoferrite assium, on solutions of copper. Crystals of the ite, nitrate, acetate, and subacetate, exhibited; also solutions. Exhibition of the protoxide, and of copper superficially dioxidized.

SECTION VI.

OF LEAD.

7. Lead is found in nature, in union with sulphur and with oxygen, newise united, in the state of an oxybase, severally with chromic, ric, phosphoric, molybdic, carbonic, and arsenic acids.

3. Lead is procured chiefly from the native sulphide, known among and mineralogists under the name of galena, which is the most abund prolific of its ores. The metal is liberated from galena by exposing fame of a reverberatory furnace, which, oxidizing and expelling the liberates the lead, partially in the state of oxide, principally in the state. The protoxide of this metal in a semivitrified state, called is largely obtained in the process of cupellation, already described means of procuring silver from argentiferous galena, or from alloys it exists in union with more oxidizable metals. (1612.)

A From any of its oxides, the metal is easily obtained by heat and

In the small way, a great majority of its combinations will yield a globule, by exposure, on charcoal, to the deoxidizing or carbo-same of the blowpipe.

1731. Properties.—The colour, lustre, and malleability of lead are well known. It fuses at about 600° F. Its specific gravity is 11.352. In large masses it is pre-eminently ductile, as it may be drawn into pipes of four inches bore; but it is too deficient in tenacity to be drawn into fine wire. It is very useful to chemists, being employed to construct the chambers and vessels used in the manufacture of sulphuric acid, of chlorine, and of the bleaching and disinfecting salts.

Of the Compounds of Lead with Oxygen.

1732. The following are the only known compounds of lead with oxygen, with equivalents of the metal, and of those oxides:—

Dioxide, or dross, probably	2 atoms lead,	208
	1 atom oxygen,	8
	,	—— 216
Protoxide,	1 atom lead,	104
•	l atom oxygen,	8
		—— 112
Bioxide,	1 atom lead,	104
•	2 atoms oxygen,	16
	,	120
Red oxide, or minium,	3 atoms lead,	312
,	4 atoms oxygen,	32
	,	344

1733. The protoxide of lead, as we find it in the shops under the appellation of litharge, is of a yellow colour when in mass, but reddish-yellow when pulverized. In the pulverulent form it is known in commerce by the name of massicot. It appears to be soluble in pure water, but is rendered insoluble by the presence of the smallest quantity of chloride of sodium or of any earthy matter. The protoxide of lead acts both as an acid and a base. In the latter capacity, it unites with the more powerful acids; in the former, with the earths and alkalies.

1734. When the protoxide of lead is powdered and heated nearly to redness, and then suffered to cool slowly, it is converted into a substance called minium or red lead, which is largely consumed as one of the materials of flint glass. Formerly minium was considered as a distinct oxide, and to this view of its composition Berzelius isclines. Thenard alleges, upon the authority of experiments made by Dumas, that it consists of three atoms of lead united to four of oxygen, and infers that it is a compound of one atom of the bioxide, and two of the protoxide

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1735. When minium is exposed to a red heat, it evolves

oxygen, and is converted into the protoxide.

1736. The bioxide of lead is obtained by the action of nitric acid on minium. If minium be a sesquioxide, one portion of it yields half of an equivalent of oxygen to another portion, and forms a bioxide, while it passes to the state of protoxide, and is taken up by the acid. But if the opinion of Thenard and Damas be correct, it must be inferred that nitric acid dissolves the protoxide, and thus extricates the bioxide previously existing in the mass.

1737. The bioxide is of a flea-colour, and is convertible by heat, first into minium, and then into protoxide. When triturated with sulphur, inflammation ensues. According to Thenard, this oxide never acts as a base, and but sel-

dom as an acid.

1738. Berzelius alleges that the gray pellicle or dross, which forms on the surface of lead when exposed to the air, and which accumulates in greater quantities when the metal is heated, is a dioxide of lead. He also states that this oxide may be obtained by the decomposition of oxalate of lead by heat. Some recent experiments of Mr. Boussin-

gault tend to confirm this opinion.

1739. The habitudes of lead with nitric, sulphuric, and chlorohydric acid, are so analogous to those of mercury with the same acids, that I do not deem it necessary to do more than point out the analogy; at the same time mentioning that, in the case of lead, no compounds are formed by oxacids with any oxide besides the protoxide; and that the resulting compounds have an insolubility more marked and invariable. Of all the important acids, only nitric and acetic acid form soluble compounds with lead. Consequently, as in any mixture, those ingredients which form insoluble combinations, always exercise a superior affinity, it follows that, from its solutions, this metal will be precipitated by any of the important salts, excepting the nitrates or acetates. Thus it will be precipitated by sulphates, phosphates, carbonates, borates, oxalates, chromates, arseniates, arsenites, tartrates, citrates, mallates, meconates, benzoates; also by any of the soluble compounds of the halogen bodies, or any of the amphydric or halohydric acids. (860.)

Of the Compounds of the Protoxide of Lead with As Acid.

and forms the acetate, called in commerce sugar of & This name was given to the acetate of lead in conseque of its taste, which is sweet and astringent. It is crysticable, soluble in water, and decomposable by heat.

1741. The acetate of lead, consisting of one atom oxide, and one atom of acetic acid, by digestion with protoxide, whether in the form of litharge or of massimay take either one or two additional atoms of oxide forming a diacetate consisting of two atoms of oxide one of acid, or a triacetate, consisting of three atom oxide and one of acid.

excess of the protoxide, a hexacetate may be obtained, a sisting of six atoms of oxide and one of acid. This c pound may be produced also, by decomposing the ace by an excess of ammonia. It forms, when dried, a w powder, slightly soluble in boiling, but insoluble in water.

1743. Goulard's extract, of which one "fluidrack agreeably to the U.S. Pharmacopæia, is to be added a pint of distilled water to make lead-water, is usually a sidered as a diacetate, and called the subacetate; but he formula, it must be evident that it may contain coordinate to make it partially, if not wholly, a triacetate.

1744. It appears to me that medical practitioners not ignorant of the difference which exists in composibetween these acetates, are too inattentive to the post

diversity of their effects.

1745. When an acetate, containing more than the portion of one atom of oxide to one of acid, is brown into contact with carbonic acid gas, a precipitate end of carbonate of lead. Hence lead-water may be used test for carbonic acid, producing results on breathing it, or upon adding it to a solution of any carbonate, and gous to those produced by lime-water under like circ stances.

Of Carbonate of Lead.

1746. When exposed to the fumes of vinegar, wi consist of acetic acid and carbonic acid gas, lead is

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by the acetic acid, and combines with the carbonic forming ceruse, or the white lead of commerce.

17. According to Thenard, the best process for obg the carbonate of lead, is to pass carbonic acid the asolution of the diacetate. Half the lead is preciding the state of carbonate, and the remainder conin solution as an acetate. The solution of acetate uced to a diacetate by boiling it with oxide of lead, abjected to carbonic acid as before. In this way care of lead of the best quality is procured, with comvely little waste of the acetic acid. Carbonate of s found in nature.

Of the Compounds of Lead with the Halogen Class.

The analogy between the habitudes of lead and mercury with acids, alluded, is not greater than that which exists between their habitudes with the haloss. Analogous reciprocal decompositions ensue, whether solutions of the xysalts of lead or mercury be mingled with solutions of chlorides, bromides, fuorides, or cyanides of the alkalifiable metals.

The chloride of lead is white, crystallizable, soluble in thirty times its weight, and has a sweet and astringent taste. When exposed to a red heat it melts, is a mass formerly called plumbum corneum, from its resemblance to horn. sat be pushed to redness, and the access of air be permitted, the chlorine is volatilized; and the remainder is found to constitute an oxychloride. Severoxychlorides exist, containing the oxide of lead, united to the chloride in proportions. One of these is found native in England.

The bromide, iodide, fluoride, and cyanide of lead, which may, as abovemente generated by means analogous to those by which the chloride is obtained, a little practical importance to make it expedient to notice them particularly. The fluoride acts as a fluobase, the cyanide as a cyanobase. The former with the fluacide of boron and silicon, the latter with cyanoferrous (fereight) acid. (1299, &c.)

Of the Compounds of Lead with Sulphur and Selenium.

Sulphur forms three compounds with lead, a disulphide, a protosulphide, resulphide. Of these, the only compound which I deem it proper to notice, otosulphide, which has already been adverted to as the principal ore of lead, lena. This sulphide may be formed artificially by heating lead and sulphur. The protosulphide of lead is tasteless, inodorous, indecomposable by heat, fusible than lead. It acts as a sulphobase, and is composed of one atom of ited to one atom of sulphur.

The selenide of lead may be procured by exposing lead, mingled with seleheat. When thus obtained it is gray, but by friction it becomes polished

be like silver. It is found in nature.

Experimental Illustrations.

4. Solution of lead in nitric acid. Its solutions preted by sulphates, chlorides, phosphates, and chro-Lead Also by sulphydric acid. Precipitation of carbonate and from the subacetate by the carbonic acid of the Lead Galena decomposed by the blowpipe flame.

SECTION VII.

OF TIN.

1755. This metal is found in the state of oxide, and in that of sulphide. The sulphide is rare, and contains much copper. The ore of tin, which is the principal source of the metal, is the bioxide which is reduced by heat and charcoal. Tin is sold in commerce under the name of block tin, to

distinguish it from tinned iron plates, vulgarly called tin.

1756. Properties.—The colour and lustre of tin may be seen in utensils newly made of tinned iron. It is very malleable and ductile; tinfol being only \(\frac{1}{1000}\)th of an inch thick. Tin tarnishes slightly by exposure to the air. It is distinguished by producing a peculiar crackling noise, when its ingots are bent to and fro. It melts at 442° F. Its specific gravity is 7.9.

1757. The equivalents of tin, and of its compounds with oxygen, chirine, and sulphur, are as follows:—

Tin -	•	•	-	•	-	59
Protoxide	. 1	atom metal,	1	oxygen,		67
Bioxide	1	"	2	"		75
Protochloride	1	>>	1	chlorine,		95
Bichloride	1	77	2	"		131
Protosulphide	1	77	1	sulphur,		75
Bisulphide	1	>>	2	"		91

Of the Compounds of Tin with Oxygen.

1758. The protoxide of tin, may be procured by adding potash to the protochleride. A reciprocal decomposition takes place between the oxide of potassium and protochloride of tin, which results in the formation of the chloride of potassium, and protoxide of tin. The former remains in solution, and the latter precipitates in the state of a white hydrate. From this hydrate the water may be expelled by heat; and a grayish black anhydrous protoxide is thus obtained, which is liable, by contact with an ignited body, to take fire, and consequently to be converted into the bioxide. The

hydrate is likewise combustible, though in a less degree.

1759. The hydrated bioxide of tin may be speedily obtained by the reaction of unfoil or tin powder with concentrated nitric acid, which is decomposed with great violence, bioxidizing the metal without dissolving it. This oxide may be obtained in the same hydrated state, by precipitation from the bichloride by an alkali. The hydrates thus obtained, though in composition the same, are different in properties. Both are soluble in alkalies, but only the latter is soluble in acids. This diversity continues even after they are severally dissolved by alkaline solutions, and subsequently precipitated by acids. These hydrated bioxides of tin are, therefore, conceived to present a case of isomerism. (1153.)

1760. We may convert the bioxide, as obtained by means of nitric acid, into the other modification, by first changing it into a chloride, and then precipitating it by potash. According to Thenard, if the precipitated bioxide be subjected to heat, it

becomes insoluble in acids.

1761. In anhydrous bioxide of tin may be obtained by subjecting tin to intense heat in contact with air. It is white, infusible, and indecomposable by heat is reddens moistened litmus paper when placed on it. This oxide is frequently found crystallized in nature. It is employed in the arts for the manufacture of enames and, under the name of putty, in grinding glass, and in making a paste for hears. The bioxide of tin acts both as an oxacid and an oxybase, combining, under favourable circumstances, with either acids or alkalies.

1762. Concentrated sulphuric acid, when cold, has no action on tin, but with the assistance of heat dissolves it, disengaging sulphurous acid gas, and forming a sulphurous acid gas, and acid gas acid

phate of the protoxide or bioxide.

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1763. The reaction above alluded to, between concentrated mitric acid and tin, when the tin is in a state of minute division, is followed by a rise of temperature, a decomposition of the acid, the evolution of nitrogen nearly pure, and the formation of the bioxide of tin. If the nitric acid be diluted so as to have a specific gravity of 1.114, and the temperature be prevented from rising by cold water, or other refrigerating applications, no gas is disengaged; since, water being decomposed simultaneously with a portion of the acid, the nitrogen and hydrogen which are thus liberated, units to form ammonia. This combines with part of the nitric acid; so that at the close of the operation we obtain nitrate of ammonia, mingled with the nitrate of the protoxide of tin.

Of the Compounds of Tin with the Halogen Class.

1764. A crystalline hydrate of the Protochloride of tin, may be obtained by subjecting the metal, in a divided state, to the action of chlorohydric acid, and then crystalling the resulting solution by evaporation. The protochloride may be procured in an anhydrous state, either by subjecting this hydrate to a temperature sufficiently high to drive off the water, or by exposing a mixture of the bichloride of mercury and metallic tin to a red-heat. When exposed, either in the solid state or in that of solution, to the action of the air, or of liquids containing oxygen, the protochloride attracts that gas, and is converted into an oxychloride. (1666.) It is probable that in this case a portion of the protochloride is decomposed, the chlorine forming a bichloride with the remainder and the liberated metal uniting with the oxygen.

1765. The protockleride of tin acts as an acid. It is composed of one atom of tin,

and one of chlorine.

1766. The bichloride of tin may be obtained in the anhydrous state, by gently heating a mixture of metallic tin with the bichloride of mercury. The bichloride is a colourless liquid, very acrid to the taste. It is volatile, and when exposed to the air produces dense and suffocating fumes. It is still occasionally called by its ancient name of the fuming liquor of Libavius. By the addition of one-third of its weight of water, this bichloride forms a solid crystallizable hydrate, which, nevertheless, dis-

solves on the addition of a sufficient quantity of water.

1767. I have been enabled to form the bichloride of tin by the direct reaction of the metal with a current of chlorine, supplied by a self-regulating reservoir. (798.) An ingot, of as much as three or four ounces in weight, was introduced into a tube of about an inch in bore, previously drawn into a capillary perforation at one end. (1372, &c.) The smaller portion of the tube was curved upwards, nearly so as to form a right angle, and being inclined towards the bend, any liquid generated within the tube, had inevitably to flow into and occupy the cavity at the curvature. By these means the operator was furnished with an index by which to regulate the supply of chlorine. The apparatus being thus constructed and arranged, the tube had, at the commencement of the process, to be filled with chlorine, and the supply of this gas afterwards so regulated as to prevent any more from reaching the included metal, than it was competent to absorb. Under these circumstances, the reaction proceeded with so much energy as actually to cause the fusion of the ingot, while an ounce measure of the bichloride was soon generated.

1768. lodine, bromine, and fluorine severally combine with tin in two proportions.

The perfluoride of tin acts as a base.

1769. The cyanides of tin have never been isolated. Berzelius, however, states, that they exist in combination with those of iron, in which case they probably play the part of cyanobases.

Of the Compounds of Tin with Sulphur and Selenium.

1770. The protosulphide of tin is obtained by heating, in a crucible, three parts of finely divided tin, and two of flowers of sulphur. This sulphide is solid, crystallizable, indecomposable by heat, less fusible than tin, and acts as a sulphobase. It is composed of an atom of tin, united to an atom of sulphur.

1771. The bisulphide of tin I have obtained by exposing to heat in a coated glass matraes, a mixture of two parts of tin, one and a half of sulphur, one of mercury, and one of sal-ammoniac." This compound, generally known as aurum musicum, or

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Bermins suggests that the mercury probably acts by bringing the tin into a state of more intimate mixture with the sulphur; and the sal-ammoniac, by carrying off, in consequence of its volatility, the heat which is evolved during the union of the sulphur with the tin, and which would otherwise be sufficient to decompose the hisuiphide, were it already formed.

mosaic gold, is of a beautiful golden yellow. When exposed to a red heat, it is decomposed. It acts feebly as a sulphobase, and powerfully as a sulphacid. This sulphide, when spread on the surface of the cushions of electrical machines, has been found to increase their exciting power.

1772. According to Berzelius there is a third sulphide, which contains a quantity of sulphur, intermediate between those which exist in the protosulphide and the

bisulphide.

1773. A gray selenide of tin may be procured by gently heating finely divided in with selenium.

Experimental Illustrations.

1774. Exhibition of tin and of tin foil; also of the funing liquor of Libavius, and the process for the generation of this chloride in which an ingot of tin is made to react with chlorine. Reaction of nitric acid or nitrate of copper with tin powder. Solution of tin by chlorohydric acid, and effects of the chloride thus obtained on some other metallic solutions. Decoloration of ink and Prussian blue. Ammonia evolved by a solution of tin in dilute nitric acid.

SECTION VIII.

OF BISMUTH.

1775. This metal is found in nature in the metallic state; usually, however, containing a little cobalt and arsenic, and sometimes sulphur. It is also found in the state of sulphide.

1776. The only ore of bismuth which is explored, is that in which it exists in the metallic state. From this it is evolved by exposure to a wood

fire, under which a hole is made to receive the melted metal.

1777. According to Berzelius, the bismuth of commerce contains iron and arsenic, and perhaps other metals. In order to purify it, it should be dissolved in nitric acid; the resulting clear solution should be mingled with water, by which a pure hydrated subnitrate of bismuth is precipitated. The precipitate being dried, is reduced by the aid of black flux* and a gentle heat.

a silvery white colour, very slightly tinged with red. It is pre-eminent is

The word flux is employed to signify a substance, which, being added to a mature which is fusible, or which contains a fusible body, promotes the fusion of the whole or a part of the aggregate. Crude flux is a mixture of nitre and cream of tartar, or crude tartrate of potash. White flux is the product obtained by designating the same mixture in a crucible, by a red-heat; whereas, when there is a double proportion of the bitartrate, an excess of carbon, causing the residue to black, it receives the corresponding designation. Black flux is resorted to, where is an object to deoxidize, as well as induce fusion. The fusion of the materals enables them more readily to move among each other, in obedience to their respective affinities, and renders it easier for metallic globules, as they are formed, is descend to the bottom of the vessel, so as to unite in one mass.

line. Thenara alleges that, when quite pure, its crystals are cubes, are so associated as to form a four-sided inverted pyramid, in which ses resemble stairs. Its specific gravity is 9.82. It is usually conlas unmalleable; yet Turner alleges that it may be hammered into while warm. Excepting mercury and tin, it is the most fusible of stals proper. Its fusing point is 476°. It is oxidized when kept in in the air; but not otherwise, unless the air be moist, in which case arnished. Bismuth combines with phosphorus, and probably in a proportion with hydrogen. Its equivalent is 71.

9. Bismuth has of late proved to be a most valuable material for estruction of thermo-electric batteries. See my Treatise on Electro-

t**ism**, page 63.

D. The reaction of sulphuric or nitric acid with bismuth, is very similat of the same acids with tim. Nitric acid, perhaps, reacts more by with the former metal than with the latter; since the addition of a quantity of concentrated nitric acid to powdered bismuth, causes the on of so much heat as to raise the temperature of the mass to red-The hydrated subnitrate of bismuth, obtained as abovementioned by ing the nitrate to water, is of a fine white colour, and has been called any of bismuth. If chlorohydric acid be present in the solution, the ate assumes the form of minute scales, of a pearly lustre, called thite. These precipitates have been used as pigments to improve aplexion, but are liable to be rendered black by sulphydric acid.

Of the Compounds of Bismuth with Oxygen.

1. There are two oxides of bismuth; one is a protoxide, consisting atom of metal, and one of oxygen; the other a sesquioxide, com-

consequently, of two atoms of metal, and three of oxygen.

Le Protoxide may be obtained by heating bismuth with access spheric oxygen, or by the calcination of the nitrate. When the sub(magistery) of bismuth is subjected successively to a caustic alkaution, and to cold water, it forms a white hydrated protoxide of
L. This oxide, when anhydrous, is yellowish, fusible at a red-heat,
of affinity for atmospheric oxygen, and easily reducible when heated

urbon or hydrogen. It acts as a base.

L. The sesquioxide of bismuth is obtained by boiling the protoxide solution of the chloride of potash or soda.* It is of a deep brown and, at a temperature a little below the boiling point of mercury, is coed. The sesquioxide of bismuth acts neither as a base nor as an

Of the Compounds of Bismuth with the Halogen Class.

Either directly or indirectly, compounds of bismuth may be produced with

salogen bodies.

In chlorine, this metal takes fire spontaneously, forming a protochloride, from the butyraceous consistency assumed in melting, received from the old sthe appellation of the butter of bismuth. This compound may also be obin the anhydrous state, by heating three parts of the bichloride of mercury of bismuth. When anhydrous, the protochloride is white, volatile, and ment: when subjected to water, a white insoluble oxychloride is formed. A crystalline hydrate of the protochloride of bismuth may be formed by distinguish in aqua regia, and evaporating the solution.

^{*} Thonard, Traité de Chimie, 6ème ed. ii. 484.

1787. Bromides of bismuth may be obtained by heating bismuth with her Iodides may be produced in like manner

1788. Fluorine and cyanogen both combine with bismuth. The cyanide, hor

is known only in a state of combination.

Of the Compounds of Bismuth with Sulphur and Selensum.

1789. Bismuth forms a bisulphide when heated with sulphur. At the me when the combination takes place, a great deal of heat is evolved. It is expetible, less fumble than bismuth, and possesses the metallic lustre and a grayma-j colour.

1790. When selenium is hested with bismuth, a crystalline selenide is form

a silvery white colour.

Experimental Illustrations.

1791. Bismuth and its oxide, exhibited. Its hue habitudes with the blowpipe, compared with those of antimony, and arsenic.

SECTION IX.

OF IRON.

1792. This metal is found abundantly in nature, principally in

with sulphur or oxygen.

different times, and in various countries. Besides these metallic times great number of stony bodies, called meteorolites, or aërolites, have in like manner. In the latter, iron always exists both in the state of oxide, and in that of metallic globules. The iron in these globules, the masses abovementioned, always contains nickel or cobalt, or Native metallic iron has also been found in small quantities, but det contain nickel or cobalt. Iron is one of the most generally distributes and the creation, and, in the state of oxide, probably the universal colouring matter.

1794. Four species of ferruginous minerals are very abundant in a magnetic oxides and sulphides, and sulphides and oxides which a

magnetic.

1795. Since ferruginous minerals, if not magnetic in the first imbecomes so by exposure to the flame of the blowpipe, the magnet is a useful test for iron. The ores of iron consist principally of the sesqui or of a compound of this oxide with the protoxide, called the black or netic oxide. The means of extricating iron from its ores, will be mean in treating of the compounds of iron with carbon, which will on the count be treated of first.

1796. Properties.—The mechanical properties of iron are too known to need description. It is the most tenacious substance in a capecially as steel, and the hardest among the malleable metals. In lity it has a still higher pre-eminence. Few metals are more easily on by the joint agency of air and moisture. In the pulverulent form, in it is reduced from the sesquioxide by means of hydrogen, iron is list become ignited by the access of atmospheric oxygen, even after it has

iron. 325

pletely refrigerated. This result is more likely to ensue, if a little aluhas been previously mixed with the oxide; since this prevents the n of the particles, and thus keeps them in that state of minute division this favourable to the success of the experiment. Iron is nearly as cult to fuse, as platinum. Its specific gravity is 7.788.

797. The equivalents of iron, and of its compounds with oxygen, chlo-

, and sulphur, are as follows:-

-	• • • • •	28
oxide,	1 atom metal, 1 atom oxygen	36
or oxide,	2 atoms ,, 3 atoms ,,	80
netic or ck oxide,	3 atoms " 4 atoms "	116
1 atom protoxide and 1	atom sesquioxide	
tochloride,	1 atom metal, 1 atom chlorine	64
quichloride,	2 atoms ,, 3 atoms ,,	164
osulphide,	1 atom ,, 1 atom sulphur	44
_l uisulphide,	2 atoms ,, 3 atoms ,,	104
lphide,	1 atom ,, 2 atoms ,,	60

Of the Compounds of Iron with Carbon, Boron, Silicon, and Phosphorus.

98. When ferruginous salts, containing carbon as a constituent, are exposed to without access of air, the iron and carbon are left in a state of combination in ous proportions. Some of these carburets, that from the oxalate, or from the tanno ate or Prussian blue, for instance, are liable to take fire when exposed to the air. '99. The process of evolving iron from its ores, comes under the fourth case of ity, in which one body in excess, combines with two others previously united. carbon with which the ore is ignited, combines both with the oxygen and metal, rerting the one into a fusible carburet, called cast iron, the other into carbonic The proportion of carbon in cast iron varies from 1 part in 25, to 1 in 15. In nerce, there are four varieties of cast iron; the white, the black, the gray, and nottled. In the white, there is the least carbon, in the black, the most; and ably, in the other kinds, less than in the black, and more than in the white kind. 00. It should, however, be understood, that cast iron is probably never a pure Usually, it contains silicon and manganese, and frequently magnesium phosphorus. This last mentioned element renders the iron less malleable at a temperature. From cast iron, the malleable metal is extricated by exposure to : and air; by which carbon, and silicon when present, are oxidized; the one g separated as a silicate of iron with the scoria, the other escaping as carbonic

O1. In some cases, malleable iron is obtained directly from the ore, by means of and charcoal.

92. Pure malleable iron is converted into steel, by being heated in contact with coal in ovens without access of air. The process is called *cementation*. By these ns, iron acquires from 1-50th to 1-120th of its weight of carbon. The bars are ered by the operation as they are seen in commerce. Broken up and welded,

form shear steel. Fused, they constitute cast steel.

According to Berzelius, the presence of manganese and phosphorus is essente the formation of good steel. Damask steel is a peculiar species, which postee the property of exhibiting waving lines on its surface, when acted on by an it is alleged by Thenard, that some experiments which have recently been its, tend to prove that this is owing to the presence of two carburets of iron; one which is blackened by the acid, while the other resists its action. I think it more that the appearance in question is owing to a mixture of iron and steel. In, however, been ascertained that a peculiar variety of this steel called wootz, hich comes from India, contains aluminium, and may be imitated by the introduction into steel, of a minute portion of that metal.

1804. A silicuret, and probably a boruret of iron, may be obtained by heating iron with a mixture of charcoal and silicic or boric acid.

1805. A phosphuret of iron is produced, when phosphate of iron is heated with lampblack. It resembles iron in colour, but is brittle, and fusible by the blowpipe.

Of the Compounds of Iron with Oxygen.

1806. Iron forms two oxides, a protoxide and a sesquioride: the former, consisting of an atom of each constituent; the latter, of two atoms of metal, and three atoms of oxy-

gen. Both these oxides act as bases.

1807. The protoxide is formed during the solution of the metal in diluted sulphuric acid. The reaction which ensues under these circumstances, is always attended by the evolution of hydrogen, arising from the decomposition of the water in combination with the acid, the oxidation of the metal, and the formation of a sulphate of the protoxide.

1808. I infer that the atom of water, which, by a union with the anhydrous acid, constitutes the aqueous sulphuric acid of Berzelius, or in other words the acid of the shops of sp. gr. 1.850, acts as an oxybase. So that the result may be ascribed to the exchange of one radical for another; an atom of iron taking the place of an atom of hydrogen. Agreeably to this view of the subject, the aqueous acid should be regarded as a sulphate of hydrogen.

1809. The protoxide of iron, forms with sulphuric acid, a green solution, which, by evaporation, yields crystals of the same colour, known in pharmacy as green vitriol, or green sulphate of iron. From a solution of this salt, the protoxide may be precipitated by an alkaline solution in the state of a white hydrate. From this hydrate the water cannot be expelled either by heat or desiccation, without causing the protoxide to acquire oxygen, either from the

water in union with it, or from the air.

1810. In consequence of this avidity for oxygen, solutions of this oxide become gradually more or less solutions of the sesquioxide; exchanging their grass green colour for that of red wine.

1811. The protoxide appears to exist in chalybeate springs, and, in its nascent state, to be soluble in water; although I do not find that other chemists are aware of the fact. Its existence in them is ascribed usually to the presence of carbonic acid; but I have observed it in the water of the Yellow Springs, which gave no precipitate with lime-water.

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1812. We have only to make a pile of silver coin, alter- (ted with disks of sheet iron, in a glass tumbler, supplied th water, in order to impart to the latter the property chalybeate spring water. In the tumbler, as in those rings, the red oxide will soon be seen precipitating, and ging, with its appropriate hue, both the liquid and the 38el.

1813. As light promotes the further oxidation and conquent precipitation of the iron, the solution of the protide, by the means which I have described, will be more rmanent in an opake vessel.

1814. There does not appear to be any mode in which

> protoxide of iron can be isolated.

1815. The sesquioxide, or peroxide of iron, also called e red oxide from its colour, which is of a dingy blood-red, ists in nature in great abundance, forming, sometimes, ge beds or masses, at other times, botryoidal, or mamlary concretions.

1816. Ochres consist of alumina, mixed with the sesquide of iron, either uncombined with water, or in the state

hydrate.

1817. The sesquioxide, as we have already stated, is ontaneously produced by the absorption of oxygen by protoxide, when exposed to the air. In fact, by the lition of nitric acid to any ferruginous solution, the iron comes more or less sesquioxidized. On the other hand, it y be partially deoxidized, and restored to the state of stoxide, by digestion with iron filings, or by the addition protochloride of tin. Hence, the black colour of the ino gallate of iron, which, when suspended in water, istitutes common writing ink, is removed by the addin of this protochloride. It appears probable, that the passes to the state of oxychloride in the following way. e portion of this metal takes chlorine from another pora to form a bichloride, while the other portion abstracts rgen from the iron, forming of course an oxide. nlting oxide combining with the bichloride, an oxychloe is produced. In the state of protoxide, to which the n is brought by the partial deprivation of oxygen, it ms a colourless compound with the tanno gallic acid.*

Protochloride of tin is the most efficient remedy for removing ink stains, or iron It is made by the reaction of chlorohydric acid with an excess of tin in der or in tinfoil, or otherwise sufficiently comminuted. It is better to use it



1818. When a solution of the protoxide of iron is added to a solution of the chloride of gold, this metal probably relinquishes its chlorine to one portion of the iron in the protoxide. The oxygen, consequently displaced, sesqueoxidizes another portion of the iron; so that metallic gold precipitates, and the chloride and oxide of iron, combining

in the state of an oxychloride, remain in solution.

1819. By intense heat the acid may be expelled either from a nitrate or sulphate of iron, and the sesquiouse consequently obtained. It has been stated, in treating of sulphuric acid, that it was originally distilled from copperas or green vitriol, the sulphate of the protoxide of iron. The oxide which remains after the expulsion of the acid, has long been known under the name of colcothar of vitriol. The metal necessarily becomes peroxidized during this process by the partial decomposition of the acid. (771.) To renrender it free from all remains of acid, it should be washed with water.

1820. The protoxide and sesquioxide of iron combine in various proportions. The scales, called finery cinder, which fly off during the forging of incandescent iron, consist of protoxide and sesquioxide. The oxide formed by subjecting iron at a red-heat to steam, is the black oxide, composed of one atom of protoxide and one of seequioxide.

1821. The native magnetic oxide of the mineralogists, is, according to Thenard, the same as that obtained when

iron is oxidized by steam.

1822. The same author alleges that neither the hydrate of the protoxide nor sesquioxide are magnetic; this quality being exhibited only when the two oxides are associated in the proportion of one atom of protoxide to one of sequioxide.

Of the Reaction of Iron with Acids.

and concentrated, is quite analogous to that already described as taking place between that acid, and mercury copper, lead, &c. The reaction of iron with this acid.

with an equal portion of acetic acid, and the addition of its volume of water. The apot to be operated upon should be first moistened with water to prevent the the ride from spreading unnecessarily. After the stain disappears, the remains of a solution should be well washed, as otherwise corresion might ensus.

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then dilute, has been mentioned and explained above.

1807.)

1824. In its habitudes with nitric acid, iron resembles in and bismuth. If the acid employed be concentrated, and the iron minutely divided, the reaction is liable to be-

ome explosive.

1825. With gallic and tannic acids, as existing in the afusion of galls, the sesquioxide of iron produces a purple r black colour, in other words, ink.* With succinic acid he sesquioxide yields a brown precipitate; with benzoic cid, an olive coloured precipitate; and with meconic and alphocyanhydric acid, a blood-red colour.

Of the Compounds of Iron with the Halogen Class.

1826. Chlorine forms with iron a protochloride and esquichloride, which correspond in composition with the xides.

1827. The anhydrous protochloride may be obtained by ussing chlorohydric acid gas over iron filings heated to

edness in a glass tube.

1828. The hydrous protochloride may be procured by the ction of liquid chlorohydric acid on iron filings. The rotochloride, in its anhydrous state, is of a pale green olour, astringent, crystallizable, very soluble in water, and olatilizable by heat. When exposed to the action of the ir, it absorbs oxygen, and forms an oxychloride, consisting of the sesquioxide, and sesquichloride.

1829. The hydrous sesquichloride of iron is produced,

The materials for common writing ink are an infusion of galls, sometimes with addition of a small proportion of an infusion of logwood, and green sulphate of m, which, in its ordinary state, contains more or less of the sesquioxide of that stal. The black hue of the liquid resulting from these infusions, increases in insity by exposure to the atmospheric oxygen, and consequent increase of the proration of sesquioxide. Dr. Ure conceives that ink made of iron, in an inferior deter of oxidizement, penetrates the paper better than that which is made by solutions the sesquioxide, and finally becomes equally black upon paper. There is some sessity respecting the composition of this liquid on account of the discordancy of insien which has existed respecting the acids of which it consists.

It may be inferred that the acid prevailing in a fresh infusion of galls, is mainly set which was formerly called tannin, and latterly tannic acid. This acid is gradully converted into gallic acid, when the infusion in which it exists is exposed to six. Either acid will produce ink, with ferruginous solutions, but it does not present to me, that it is known which of the two answers the best for this purpose. have found a beautiful blue black ink to result from the reaction of a filtered insien of galls in cold water with finery cinder. It is too much prone to precipitate, at by agretion is always resuspended. The old practice of introducing cotton into a inhatend, removes this inconvenience in great measure. Over common ink it set these advantages, it contains no free sulphuric acid, and makes no grounds which

when the sesquioxide of iron is exposed to the action of chlorohydric acid. It may be obtained in the anhydress state, by heating iron filings in an excess of chlorine. Thus obtained, it is volatile and deliquescent.

1830. Bromine and iodine form compounds with iron, which no doubt correspond in composition with its oxides and chlorides.

1831. There are two fluorides of iron which : it either as acids or bases.

1832. The protocyanide of iron is formed by exposing the cyanoferrite of annual nium, which is a compound of the cyanides of iron and ammonium, to heat it a start tort. The cyanide of ammonium, which is vertile, passes over, leaving the protocyanide of iron in the form of a grayish-yellow powder. This cyanide all as a powerful cyanacid, combining in that capacity with the cyanides of almost at the metals. It also combines with cyanhydric acid, but whether as an acid or a tass, appears to me doubtful. I incline to the opinion that it acts as an acid, faming a cyanoferrite of the cyanobase of hydrogen.

1833. The sesquicyanide of iron is obtained by mingling a solution of the familicate of the fluobase of iron with a solution of the cyanoferrite of potassium. A familicate of the fluobase of potassium precipitates, and the sesquicyanide of iron mains dissolved. Its solution is of a deep brownish-yellow colour, and an astringuate. If we attempt to obtain it in the solid form by desiccation, it is pastily

decomposed, and converted into Prussian blue.

1834. Of Pressian Blue.—When the cyanide of potassium is mingled in stable with a ferruginous salt, a precipitate ensues, well known under the name of Paintellius, having been first accidentally discovered at Berlin. It would seem, that perfect the colour of this precipitate, both oxides of iron should be present; in the protoxide may produce the protocyanide, and the sesquioxide the sesquioxide These cyanides, by their union, form the compound in question. (1299, &c.)

Of the Compounds of Iron with Sulphur and Selenium.

1885. Iron forms with sulphur a protoculphide, a sesquisulphide, and a histiphile.

Moreover, the protoculphide combines in various proportions with the histiphile.

with the metal.

1836. Hydrated protosulphide is alleged to be formed during the combustion which arises from triturating with moisture two parts of iron filings with one and a initial sulphur. This hydrated protosulphide is liable to absorb oxygen with a rapidity so great as to produce ignition. Owing to this property, its presence in hiteminous coal beds sometimes causes them to take fire spontaneously.

1837. Native protosulphide of iron is of rare occurrence; but the magnetic and bisulphides are abundantly found in nature, especially the latter, which is one of the most common minerals. From its resemblance to gold, it is frequently mistakes for that metal by inexperienced observers. When intensely heated, a portion of its subphur sublimes; and hence it is one of the sources of that important substance.

1838. Of the bisulphide, it is alleged by Thenard, that there are two variations which, though identical in composition, are dissimilar in their crystalline forms in their properties. Of these varieties, only one is susceptible of spontaneous restion with air and moisture, and consequent conversion into a sulphate. To a similar transformation of this and other sulphides, we are indebted for the greater part of the green vitriol, or sulphate of iron, used in the arts. Beds of these minerals, in a state of decomposition, are to be met with in every country.

1839. Sesquisulphide of iron is produced, when the sesquioxide of this metal is exposed to a current of sulphydric acid, provided the temperature be not above as

At a higher temperature, a bisulphide results.

1840. The protosulphide and bisulphide of iron, constitute, as Thenard mentions, the mineral called magnetic pyrites. This mineral is also formed, as he alleges, when iron in a state of intense ignition is presented to sulphur, and when either the sequisulphide or bisulphide is fused. In fact, it would seem that he considers need the other sulphides as magnetic; although the presence of a greater proportion iron in the protosulphide would lead us to suspect in it a greater susceptibility of magnetic influence. Berzelius, however, considers the protosulphide as magnetic.

1841. The selenide of iron is formed by causing the vapour of selenium to per over iron filings heated in a glass tube. It has a metallic brilliancy, and a deep gray

colour approaching to yellow.

Experimental Illustrations.

12. Iron, dissolved by chlorohydric and sulphuric Red and magnetic oxide of iron, exhibited; and solutions precipitated by galls, and by cyanoferrite assium. Effects of protochloride of tin on the colour; precipitates. Ores of iron, rendered magnetic by owpipe.

SECTION X.

OF ZINC.

L This metal exists in nature in four states; in that of sulphate, silirbonate, and sulphide. As a silicate or carbonate, it is known in ogy under the name of calamine; its sulphide is called blende.

From calamine or from blende, when converted into an oxide by the metal is obtained by heating it with charcoal, in a crucible tole in the centre of the bottom. To this a sheet iron tube is adapted the sinc is conveyed in liquid globules or vapour to a vessel of situated beneath, within which the vapour consequently condenses, occas is called distillation by descent, "distillatio per descensum."

ay be purified by redistillation.

• Properties.—Zinc is of a brilliant metallic white colour, tinged has of lead. Its structure is strikingly crystalline. Its specific is about 6.86. Under ordinary circumstances it is not malleable, be laminated by rollers at a heat somewhat above that of hoiling It melts at about 680°. That it may be volatilized at a higher temmust be evident from the process by which it is obtained as aboved. (1844.) By exposure to the atmosphere it is slightly oxidized, white heat burns rapidly with intense light, the resulting oxide betilized in fumes. Water is rapidly decomposed when passed in the steam over ignited zinc, or when presented to it together with a due on of sulphuric or chlorohydric acid. Zinc combines with carbon sphorus.

The equivalents of zinc, and of its compounds with oxygen, chlo-

i sulphur, are as follows:—

ıc, · ·	•		•		• ,	32
toxide,	1	atom	metal,	1 oxygen,		40
oxide, doubtful.	1	• ••	22	1 chlorine,		68
phide,	1	• • • • • • • • • • • • • • • • • • • •	"	1 sulphur,		48

Of the Compounds of Zinc with Oxygen.

7. The protoxide of zinc is formed during the comn of the metal in atmospheric air. From the lightness and fleeciness of its texture, when obtained in this way, it was formerly variously called pompholix, nihil album, or lana philosophica. The protoxide may be obtained from one variety of the ore called calamine, by heating it to expel carbonic acid. To prepare it as it is presented to us in the shops, the ore is roasted, pulverized, and levigated. A better process, as I conceive, is that of collecting the woolly matter produced by the combustion of the metal. But to either of these modes I should prefer that of precipitating the oxide from the sulphate in solution, by liquid ammonia.

1848. Peroxide of zinc has been obtained by mingling bioxide of hydrogen with a dilute solution of the nitrate of this metal, as in the process for the peroxide of copper, which it resembles in many of its properties. (1706.) The protoxide usually acts as a base, though in some cases it may act feebly as an acid. The peroxide performs the

part neither of a base nor of an acid.

1849. The reaction of sulphuric acid with zinc is similar to that of the same acid with iron. (1807.) When subjected to nitric acid, zinc takes all the oxygen from one portion of the acid, while the protoxide thus formed is dissolved by another portion; meanwhile the nitrogen escapes with violent effervescence. Professor Emmet has recommended the reaction of this metal with the nitric acid in nitrate of

ammonia, as the means of procuring pure nitrogen.

1850. If the solution of the acetate of zinc, obtained by the reciprocal decomposition of the acetate of lead and sulphate of zinc, (522,) be clarified by subsidence or filtration, and then evaporated, the acetate of zinc may be obtained in the crystalline form. It will also be in a state of purity if the materials have been used in the equivalent proportions, or with a slight excess of the acetate of lead-Pure acetate of zinc may also be obtained by the process for forming the arbor Saturni; as in that process, after a sufficient time, the lead is completely precipitated by the zinc, which remains in solution. In this process a piece of zinc being suspended in a solution of acetate, or, preferably, nitrate of lead, and having a greater affinity for oxygen, it deoxidizes the lead. This, being thus rendered insoluble, precipitates; while the resulting oxide of zinc is seized by the acid and dissolved.

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o one of the formulas of the Pharmacopæias, as a tincure, in other words, in alcoholic solution, by subjecting a nixture of sulphate of zinc and acetate of potash, in equivalent proportions, to alcohol. The mixture of the salts sollowed by a reciprocal decomposition, analogous to that produced by the mixture of sulphate of zinc, and acetate of lead; excepting that the resulting sulphate of lead is quite insoluble in water, and separates by precipitation; while, in the other case, both of the resulting salts, being more or less soluble in water, alcohol is employed to separate them. This liquid does not dissolve the sulphate of potash, while it readily takes up the acetate of zinc.

Of the Compounds of Zinc with the Halogen Class.

1852. Anhydrous chloride of zinc is formed during the combustion of zinc in chlorine. It was formerly called the butter of zinc, from its consistency. It is of a grayish-white colour, translucid, astringent, fusible at the temperature of boiling water, and volatilizable at a red-heat. By dissilving zinc flings in chlorohydric acid, and evaporating the solution to drypess, we may obtain this chloride in the state of hydrate.

1853. Zinc combines with iodine, fluorine, and cyanogen. The cyanide

acts as an acid, the fluoride both as a base and an acid.

Of the Compounds of Zinc with Sulphur and Selenium.

1854. Sulphide of zinc may be obtained by heating the sulphate to whiteness with a carbonaceous paste. It is difficult to combine zinc directly with sulphur; but when the vapour of sulphur is passed over incandescent zinc, a combination takes place with a violent commotion, and the evolution of so much heat as to volatilize part of the zinc. The same result ensues when zinc filings are suddenly and intensely heated with the persulphide of potassium, or the powdered bisulphide of necessary.

1855. Sulphide of zinc is solid, yellow, tasteless, less fusible than zinc, indecomresable by heat alone, but reducible by intense ignition with charcoal. It is a power-

ul sulphobase.

1856. When the sulphate of zinc is decomposed at a low red-heat by hydrogen, a expenience, or in other words a compound of the sulphide and oxide, is formed.

1857. When the vapour of selenium is passed over zinc heated to redness, the miss of the two substances takes place with violence, being attended with the phenomena of active combustion. The resulting selenide is a yellow powder.

Experimental Illustrations.

1858. Zinc, subjected to diluted sulphuric, and diluted chlorohydric acid. Arbor Saturni, produced by it in a solution of nitrate of lead. Combustion of the metal in an incandescent crucible. Its habitudes with the blowpipe,

exhibited. Reaction of zinc filings and bisulphide of mercury; also of the melted metal with a fused nitrate.

SECTION XI.

OF ARSENIC.

1859. This metal is found in nature, in combination with oxygen, sulphur, and various metals. It is sold in commerce under the name of cobalt, and in the state in which it bears this name, it is full of crevices, and so much tarnished or blackened by oxidizement, both internally and externally, that it is not possible, even by a fresh fracture,

to see the true colour and lustre of the metal.

absurdly named) should be coarsely pulverized, and introduced into a glass tube sealed at one end. The tube should be less than half full. Thus prepared, it should be placed within a cylinder of iron, closed at the base. The butt-end of a gun barrel will answer. The space between the iron and the glass should be filled with sand, and another gun barrel applied, so as to receive any fumes which may arise, and conduct them into a chimney. That portion of the glass tube which contains the arsenic, should be kept red-hot for about half an hour. After the apparatus is quite cool, the metal will be found in crystals of great splendour, occupying that portion of the glass tube which is next the part heated to redness.

1861. Properties.—Exposed before the blowpipe, arsenic is distinguished by burning before it fuses, and by emitting copious white fumes, which have the odour of garlic. These fumes are easily produced, by projecting a portion of the metal upon a hot iron, or by subjecting it in any other way to heat and air. They are evolved on a large scale during some metallurgic operations, and, after being purified by a subsequent sublimation, constitute the arsenious acid or white arsenic of the shops. This metal is extremely brittle and friable, and, when newly sublimed, has the colour and brilliancy of polished steel. It requires less heat to vaporize than to fuse it; so that it cannot be melted without the aid of a pressure greater than that of

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tort filled with nitrogen, at the temperature of 356°. mder the impression, that the nitrogen must coas a solvent in this result; taking up the metal in rmer part of the retort, and depositing it in the part. I have ascertained that metallic arsenic, I in a glass tube immersed in melted lead, is not zed, unless so far as it may be oxidized; and, morethe process for obtaining the arsenical ring, I have ed that it is formed just beyond the part of the tube s reddened by the heat.

2. The following table gives the equivalents of , and of its compounds with oxygen, chlorine, and

,		-		38
le, doubts	ul.			
ous acid,	2 atoms	metal, 3	atoms oxygen,	100
: acid,	2 atoms	,, 5	atoms "	116
ıloride,	1 atom	,, 1	atom chlorine,	74
:hloride,	2 atoms	,,	atoms "	184
ılphide,	1 atom	,, 1	atom sulphur,	54
rulphide,	2 atoms	_	atoms ,	124
hide,	2 atoms	,, 5	atoms ,	156

Of the Compounds of Arsenic with Oxygen.

. According to Berzelius, the black matter which s the brilliancy of metallic arsenic on exposure to , is a suboxide. Thenard seems inclined to conas a protoxide; while by other chemists it is treat-. mixture of arsenious acid and the metal; as, when I to heat or to acids, it yields arsenious acid, and c arsenic. But as it appears that arsenious acid is ound of oxygen and the metal, in the ratio of three of the former to two of the latter, it would be reato infer the existence of a compound consisting of m of each. Besides, it has been ascertained by us, that the exposure of arsenic to air never causes orption of more than eight per cent. of oxygen; s, to form arsenious acid, the metal must absorb wo per cent. Now it seems very improbable that, he same circumstances, one portion of the metal

should absorb thirty-two per cent. of oxygen, while ano-

ther portion should absorb none.

1864. Arsenious acid is found in nature both in crystalline form and in that of white powder. It forms the fumes which are so copiously evolved when arsenic is ignited in the air. It is milk-white, has a rough and slightly acid taste, followed by a flavour feebly sweet. It is hardly necessary to state that it is a virulent poison. When subjected in open vessels to a low red-heat, it softens, and sublimes, in the form of a white powder, or, when the vessels are large and the operation slow, in regular octobedral crystals.

1865. Arsenious acid is soluble in water, but not to any great extent. Berzelius states that a saturated solution of it in boiling water, in which the deposition of crystals has commenced, contains but a twelfth or thirteenth of its weight. There is much uncertainty, and some mystery, respecting the extent of its solubility in cold water. Berzelius quotes an observation made by Fischer, that the portion employed is never entirely dissolved; and that as the ratio of the water to the acid increases, this being always in excess, the quantity dissolved lessens. Thus 80 parts of the former take up ½0th of its weight; 160 parts, ½0th; 240 parts, ½10th; and 1000 parts, only ½200th.

a heat approaching to redness, fuses into a transparent glass of the specific gravity of 3.699, unchangeable in dry air, but gradually becoming white and opake in a humid atmosphere. In the last mentioned state, it appears to be more soluble in boiling water, and to be retained in solution to a greater extent than in the transparent state. The transparent acid reddens litmus; while, by the opake, litmus previously reddened may be restored to its original colour. These varieties of arsenious acid are, therefore.

considered as isomeric. (1153.)

1867. Of Arsenic acid.—By digestion in aqua regia or in strong nitric acid, evaporation of the resulting solution to dryness, and subsequent ignition nearly to redness in a platinum crucible, arsenious acid acquires two additional atoms of oxygen; so that a compound is formed in which the metal is to the oxygen, in the proportion of two atoms to five. This compound is arsenic acid, which is solid white, and caustic, and capable of reddening litmus.

When exposed to heat it melts into a glass; but if the teat be pushed to redness, it is decomposed into arsenious acid and oxygen gas. This is a more powerful acid, a nore virulent poison, and more energetic in its affinities, han arsenious acid. Like other acids, which bear a high emperature without decomposition or volatilization, it expels, when aided by heat, the volatile acids from their combinations. It forms, with certain metallic oxides, salts which crystallize in the same form as the corresponding phosphates; whence, as I have elsewhere stated, (474,) arsenic and phosphoric acid are said to be isomorphous. Of such bodies, one may be substituted for the other in crystalline compounds, without altering the form of the resulting crystals.

1868. Arsenic acid is deliquescent, and much more soluble than arsenious acid; yet after being vitrified by heat, it does not dissolve completely at first, but deposites a white powder, which, by frequent stirring, finally dissolves. In consequence of this and some other differences in their properties, it has been supposed that the melted and unnelted arsenic acids are isomeric with regard to each

other.

1869. Arsenious and arsenic acid severally combine with the metallic oxides. Arseniate of potash is formed, when arsenious acid or metallic arsenic is deflagrated with aitrate of potash. Fowler's solution, the liquor potassæ arsenitis of the U.S. Pharmacopæia, is made by boiling arsenious acid and carbonate of potash, of each 64 grains, with a pint of distilled water, and adding four fluidrachms of the spirit of lavender. The arsenious acid, displacing the carbonic acid, forms with the alkali an arsenite of potash. This solution produces a yellow precipitate with nirate of silver, without the aid of ammonia, as the place of his base is supplied by the potash.

1870. The soluble arsenites and arseniates yield precinitates with solutions of copper and silver, and destroy the
nue colour of the iodide of starch, by the superior affinity
of iodine for arsenic. In the instance of copper and silrer, an arsenite or arseniate of those metals is formed.
The arsenite of copper is of an apple-green colour, and forms
or pigment called Scheele's green. The arsenite of silver is

rellow; the arseniate, brick-red.

1871. Sulphuric acid when cold does not react with ar-

senic; but when warm, the acid is decomposed, and armious acid formed.

1872. Of nitric acid the reaction with arsenic is similar to the reaction of sulphuric acid with the same metal; except that it takes place without the aid of heat, and that the arsenious acid which is at first produced, is finally converted into arsenic acid.

Of the Compounds of Arsenic with the Halogen Class.

1873. A sesquichloride of arsenic is obtained by the direct reaction of chlorine with arsenic, or by the distillation of this metal with the bichloride of mercury. If, in this process, the protochloride of mercury be substituted for the other, a protochloride of arsenic is generated: and by the reaction of the metal with an excess of chlorine, a perchloride results.

1874. The sesquichloride is a colourless, fuming liquid, of an oleaging consistency, quite analogous, both as to the means of evolution and its perties, to the bichloride of tin, or fuming liquor of Libavius. (1766.)

1875. Bromine and iodine severally form compounds with arsenic, which

correspond in composition with the sesquichloride.

1876. The fluoride of arsenic is a colourless, furning liquid, which published consists of two atoms of arsenic, and three of fluorine.

Of the Compounds of Arsenic with Sulphur and Selenium.

1877. There is scarcely any limit to the number of proportions in which arsenic and sulphur appear to be capable of combining; yet Bernelin at mits the existence of but five distinct sulphides, and Thenard recognisms only three,—a protosulphide, a sesquisulphide, and a persulphide. By the union of these with various quantities of the metal, or of sulphur, all the other compounds are supposed to be produced.

1878. The proto, sesqui, and persulphide severally combine with sulpho-

bases, as sulphacids.

1879. The protosulphide of arsenic, known in commerce by the name of realgar, may be obtained by heating a mixture of two parts of sulphur, and rather less than three and a half parts of arsenic. It is procured in the large way by distilling arsenious acid with sulphur. It is tasteless, crystallizable, less fusible than arsenic, and of an orange-red colour. When heated in close vessels it volatilizes unchanged, but if the air be admitted it is converted into arsenious and sulphurous acid. It is found native.

1880. The sesquisulphide is obtained by adding chlorohydric acid to a mixed solution of sulphide of potassium and arsenite of potash. The oxygen of the arsenious acid and of the potash unites with the hydrogen of the chlorohydric acid, the chlorine with the potassium, and the sulphur with the arsenic. The chloride of potassium remains in solution, while the arsenic and sulphur precipitate in the state of sesquisulphide, and in the form of beautiful yellow flocks.

1881. The sesquisulphide is found in nature, and is known in commerce under the name of orpiment. It is crystallizable. When heated gently is close vessels it melts, and if the heat be further elevated, volatilizes, and may be condensed unchanged. If the access of air be permitted during the exercises sulphyrous and accessions will any formed.

the operation, sulphurous and arsenious acid are formed.

1882. The persulphide of arsenic is formed by passing sulphydric acid

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h a solution of arsenic acid in water. It is yellow, and resement. It is fusible, volatilizable, and capable of reddening litmus. It is fusible, volatilizable, and capable of reddening litmus. It is fusible, volatilizable, and capable of reddening litmus. It is fusible, when arsenic is dropped into selenium, liquefied by heat. If this selenide be subjected to distillation at a perselenide is obtained.

e Compounds of Arsenic with Phosphorus and Hydrogen.

I phosphuret of arsenic may be formed by heating phosphorus et al. It is black and brilliant.

or the evolution of hydrogen, (797, &c.) an aqueous solution of cid be substituted for water, the other materials being as usual cid and zinc, arseniuretted hydrogen will be evolved with no less in that with which the evolution of the pure gas is accomplished, reenious acid is not present.

Then this gas is made to pass through a tube kept by means of a coal fire, as hot as the glass will bear, the arsenic is precipie metallic form, in the cooler part of the tube, just beyond the

'he process above described, is decidedly preserable to any other, respects convenience and economy, but the safety of the ope-

'o procure this gas devoid of pure kydrogen, Soubieran recoman alloy of equal weights of arsenic with zinc be made by

subjected to strong chlorohydric acid.

confidence is to be placed in the recommendation of the distinemist above named, it follows, that by the introduction of such to the self-regulating reservoir, substituting strong chlorohydric sulphuric acid, there would be a supply of pure arseniuretted t command.

rseniuretted hydrogen is highly inflammable, in common with riform compounds of hydrogen. It is extremely deleterious to njurious when liberated in quantities too small to be immediately the operator. It is productive of nausea and vomiting, sometimes of purging. As palliatives of these symptius recommends warm tea, and sulphydric acid gas.

generating this gas for illustration, by the apparatus employed peophical candle, (805,) I inadvertently inhaled enough to prosient indisposition. Gehlen, a respectable German chemist,

by a similar inadvertency. This gas is the more insidious, a not warned of its presence by the fetidity of its odour, as in the combinations of hydrogen with sulphur and phosphorus, and nees. A ten-thousandth part of this gas may be detected in a ture, by the metallic pellicle which it causes upon a solution of blimate.

l of turpentine appears to form a crystalline compound, by re-

arseniuretted hydrogen.

solid compound of arsenic with hydrogen has been made, by n alloy of potassium and arsenic to water; and likewise by the on of water by the Voltaic series, one of the wires, employed for terminating in a piece of arsenic immersed in that liquid.

Experimental Illustrations.

1894. Appearance and habitudes of arsenic, in its metallic and crystalline form, contrasted with those of zinc, antimony, and bismuth. Arsenious acid and its solutions, exhibited; also, Fowler's solution, or solution of arsenite of potash. Arsenious and arsenic acid, in solution, added to large vessels of clear water, and detected by sulphydric acid, or by ammoniacal nitrate of silver or copper. Same acids precipitated by lime-water. Exhibition of Scheele's green, or arsenite of copper. Combustion of arseniuretted hydrogen, displayed.

Of the Means of detecting Arsenic, in Cases where poisoning by Arsenical Compounds is suspected.

1895. As respects arsenic, the most important object of attention is the means of detecting this metal, in cases in which an arsenical compound may be used as a poison.

1896. The first steps are of course directed to the collection and preservation of all the matter which may have come from the patient in vomiting, or which may be obtained by opening the stomach. As the combination of this metallic poison usually administered is arsenious and in the pulverulent form, all the matter collected and the surface of the stomach should be rigidly examined, in order to detect any particles which may remain in that state. Berzelius counsels us especially to scrutinize those spots to ascertain whether any particles of the poison are lodged in them.

1897. In the next place, the whole mass, collected and preserved as above advised, should be thrown into water, which, while stirred to cause the suspension in it of the lighter portions of the matter, should be poured off, together with that lighter matter from the heavier subsiding portion. The liquid thus separated should be filtered; and both the resulting filtered solution, and the heavier matter which may have sunk to the bottom of the vessel, should be evaporated to dryness in an appropriate oven, or in a vessel kept hot by boiling water. The whole being quite

i, it should be introduced into a glass or porcelain ves-; and, adding a sufficient quantity of strong nitric acid cover the mass, it should be subjected to a heat adequate cause a brisk reaction. This should, if necessary, be tained by further additions of the acid, until there is no

ger any organic matter undecomposed. 1898. As nitric acid can have no other effect upon aric than that of converting it into arsenic acid, in which te it is less volatile than in any other; this process tends the same time to annihilate the organic impurities, and secure the metal. Thus the matter to be assayed is ich diminished in bulk, and if it contain arsenic must ld it as arsenic acid, which is more soluble than arsenis acid. Hence, if the dry mass be digested with water, solution will be obtained, which, being filtered, may be cipitated by lime-water. The arseniate of lime which cipitates, being dried, should be mingled with about :-fourth of its weight of powdered charcoal, and introced into a glass tube, sealed at one end. The mixture ving been made to settle down to the sealed end within narrow limits as possible, a little cotton wick must be stened to the end of a wire, by twisting the wire so as to rm one end into an eye, and passing the wick through eye, and winding it about the end of the wire, until a ig of cotton be made just large enough to slide in the e like a piston. For greater security, the wick may be and about the wire, so that one end may be held in the By means of the piston, thus formed, the portion the tube, not occupied by the mixture, may be wiped ean. The tube should now be subjected to the flame of spirit lamp, the piston being retained in it, as near the ixture as it can be without being injured by the heat. ne heat should be applied at first to the anterior part of e mass, proceeding to the posterior part afterwards; and soon as the whole ceases to give out aqueous vapour, e piston should be passed quickly down, so as to wipe way the moisture condensed, together with any accomunying foulness. The piston being again beyond the ach of the heat, the part of the tube containing the mixre should, by the aid of a blowpipe, be exposed to a temrature as high as the glass will bear. If there be arsenic the mixture, it will now appear in a bright metallic ring, st beyond that part of the tube which was heated redhot. On cutting the tube at the part where the ring appears, and heating it by a spirit lamp, the alliaceous smell of arsenic will be perceived, if this metal be present; and the same smell will be experienced on igniting the cotton of

the piston above mentioned.

1899. The process thus described for obtaining the arsenical ring is nearly the same as that which I employed in the analysis of some matter sent to me from Westchester by Dr. Thomas; having been obtained from the stomach of a woman poisoned by her husband. I afterwards repeated it successfully at Westchester, in presence of Dr. Thomas and another physician; and upon their cridence of the result so obtained, the murderer was convicted. Before his execution he confessed himself to be guilty.

of the stomach, it may be found in the membranes, or coats. Hence, in making the examination for arsenic, the stomach should be boiled in nitric acid, until all the organic

matter is destroyed.

1901. Very minute quantities of arsenic may be detected by the aid of silver or copper in solution. With silver, arsenic acid gives a brick-red, arsenious acid, a yellow precipitate. With copper, arsenious acid produces a very striking green precipitate of arsenite of copper, called Scheele's green. Sulphydric acid gas produces, with either acid, a yellow precipitate of sulphide of arsenic. As these results are precarious, and hable to be produced by other causes, they should not be considered as conclusive evidence.

1902. In the case of the murder above mentioned, I found that the arsenic acid, as procured from the contents of the stomach, would not assume the appropriate hue in precipitating with silver, whether before or after its union with lime. Instead of a brick-red, it was of a muddy colour. By Dr. Feutchwanger, who was present during many of my experiments, this was ascribed, correctly, as I believe, to phosphoric acid. It has been stated, that it is difficult to separate these acids when associated, from their isomorphism, or, in other words, crystallizing in the same form. (471.)

acid can be precipitated by the soluble salts of silver or

343

- er. Hence an alkali must be present, either in union the arsenical acids, or with the metallic salt. This t is attained conveniently, by the addition of ammonia entrate of silver or copper; as, with either of those s, that alkali forms a soluble ammoniacal nitrate.
- 14. A great improvement in our means of detecting ic has been introduced by Marsh, of London. It has mentioned, (1885,) that if in the process for evolving gen by a self-regulating reservoir, an aqueous solular arsenious acid be substituted for pure water, the ials and manipulation being otherwise the same, in of pure hydrogen, arseniuretted hydrogen will be geed. From the observations of the ingenious mechaabove named, it appears, that if any mixture cong arsenic be added to water and acid, used in an gous miniature apparatus, the nascent hydrogen will ne with the arsenic, however minute the proportion. equently, a jet of the gas when inflamed, by its hue, and odour, and still more when allowed to play upon urface of a piece of porcelain or glass, will demonthe presence of arsenic. In this last mentioned case, k stain will be made, consisting of concentric circles, ich that which is central will have the metallic hue of rsenical ring, especially when examined through the
- 15. The great objection to this process, as it came the hands of Marsh is, that when the proportion of ic is very small, the whole may escape before the tor may be enabled to detect it. Hence, it would preferable to resort to the expedient recommended ubieran, of passing it through a tube heated by a (1886); or to pass all the gas generated through liquid competent to effect a complete absorption of senic. For this purpose a solution of corrosive submight answer. (1891.)

Experimental Illustrations.

6. Small portions of arsenious acid, or of the are of lime, mingled with powdered charcoal, and subto heat in a glass tube. Arsenical ring, produced chibited.

with an aqueous solution of arsenious acid, sulphuric acid, and zinc. (796, &c.) Deposition produced by the flame upon glass, mica, or porcelain. Gas passed through a glass tube, reddened by a lamp, or gas flame, deposits metallic arsenic in a film resembling in appearance the arsenical ring.

SECTION XII.

OF ANTIMONY.

1908. Antimony sometimes occurs in nature in the metallic state and a that of oxide, also abundantly as a sulphide. It is in that to the sulphide that the name of antimony is given in commerce, the metal being designated as the regulus of antimony.

1909. The ores of this metal had been known for a long time; but for its extraction from them, the world is indebted to Basil Valentine, will lived towards the close of the fifteenth century. Since that period, families utility in medicine, it has been eminently an object of investigation.

its utility in medicine, it has been eminently an object of investigation.

1910. Metallic antimony may be obtained by roungling the suphide with two-thirds of its weight of bitartrate of potash, and one-third of its weight of intre, and deflagrating the mixture in a red-hot crucible. The origin of the intre converts the sulphur into sulphurous acid, which uscapes: which the alkali of both the salts operates as a flux, or in other worth promotes the fusion of the mass. The carbon of the tartaric acid counteracts the oxidizement of the metal.

1911. Charcoal, intimately intermingled with carbonate of potash or

soda, may be used instead of the bitartrate.

1912. Antimony thus obtained is not quite pure. To render it so it may be dissolved in aqua regia, precipitated in the state of oxychloride by water.

and revived by ignition with bitartrate of potash-

1913. Properties.—Antimony is so brittle as to be easily pulserized by the processorted to in the case of sulphur. (750.) When quite pure and newly fractured, it is of a silver-white colour, and very brilliant. If it be rubbed between the fingers, they acquire a perceptible odobr. Its specific grapty is 6.7. It fuses a little below a red-heat. When thrown in a state of fusion upon a board, it produces a beautiful effect, being dispersed into a multitude of ignited globules, which emit copious times of oxide, and team their traces upon the board. The temperature of the globules seems to be supported by their own combustion.

1914. A single globule of the metal, being brought to a state of ignation by the blowpape flame upon a piece of charcoal, if held, after operation of the blowpape is descontinued, in a current of air, such as exists assistly at an aperture in a flue, will, in consequence of the heat arising from its uncertainty that the atmospheric oxygen, continue at a bright red-heat until meanity

consumed.

1915. According to Berzelius, the purity of antimony is indicated by a

whiteness, and a granular or fine lamellar texture; whereas the metal ise does not excel tin in whiteness, and is coarsely lamellar, almost asceptible of cleavage. It appears to me, that the differences here reto, are dependent, as in other cases, on slowness or quickness of coolabutton which was granular when taken from a crucible refrigerated ex,—by fusion in an iron mortar in which it was prevented from coolackly by proximity to a fire, acquired a lamellar texture. Bad anti-ooks like hornblende rock when broken, as to its crystalline texture, nnot be fused into a globule as liquid or pure. It will not answer as r the experiment of throwing on the board, as the pure metal.

3. The equivalents of antimony, and of its compounds with oxygen,

e, and sulphur, are as follows:—

ny, -	-	•	•	•	•	•	-	•	•	64
oxide,		2	atoms	metal,	3	oxygen,		-	•	152
onious acid,		2	>>)	4			-	•	160
onic acid,		2))	? ?	5			•	•	168
chloride,		2	22	77	3	chlorine,		•	•	236
ride,		2	3 7	"	4	22		-	•	272
oride,		2	22	22	5	? ?		•	•	308
ulphide,		2))))	3	sulphur,		•	•	176
ide,		2	"	"	4	27		•	-	192
hide,		2	"	>>	5	33 ·		•	•	208

Of the Sesquioxide of Antimony.

17. Sesquioxide of antimony may be obtained by exg the metal to heat with access of air; by moderately ing the sulphide; or by subjecting the sesquichloride ater, in which case a powder precipitates, called r of Algaroth, from the name of the physician who ecommended it to public attention. This powder, an oxychloride, by digestion with the carbonate of h, is converted into the sesquioxide. It may also be ned by the reaction of the metal with diluted nitric afterwards repeatedly digesting the resulting subsalt ter, until this liquid no longer reddens litmus. rm in which the sesquioxide is obtained by heat and received formerly the name of argentine flowers of ony. When obtained from the oxychloride, the seside has a tinge of gray. If the sesquioxide, as proby the last mentioned method, be heated, it takes nd is converted into antimonious acid.

18. Sulphuric acid, when cold or diluted, does not retth antimony, but, when warm and concentrated, is the decomposed, evolving sulphurous acid, and form-sesquioxide of the metal, with which the undecomacid combines. By water the acid may be for the

most part removed from this sulphate, so as to cause in an excess of oxide so great as to render it competent; the production of tartar emetic, by digestion with the tartrate of potash. In this case, the excess of oxide in a sulphate, and the excess of acid in the bitartrate, usi converting the latter salt into the double tartrate of p ash and antimony, or tartar emetic.

1919. The sesquioxide acts feebly both as an acid and base. Combined with bitartrate of potash, it constitute tartar emetic, and is the only compound of antimony we

oxygen, which is considered as medicinal.*

1920. Tartar emetic may be considered as consisting

Two equivalents of tartaric acid, One of sesquioxide of antimony, One of potash, Two of water,

 $9 \times 2 =$

 $66 \times 2 = 1$

Of the Compounds of Antimony with Oxygen, of inferior imports medicinally.

1921. Antimonious acid is generated by digesting antimony in a acid, evaporating the liquid to dryness, and calcining the residue; or thoroughly roasting the sulphide of antimony with access of air. And nious acid is white, tasteless, infusible, fixed, indecomposable by best, insoluble in water.

1922. When nitric acid is added to a solution of the antimonial potash, the antimonious acid is precipitated in the state of hydras. Is the

state it reddens litmus paper.

1923. Anhydrous antimonic acid is obtained by subjecting the cayed ride to the action of nitric acid, and afterwards exposing the resulting to a temperature of 500° or 600°, to expel any excess of this acid. By fingrating the metal with four times its weight of nitre, dissolving in the resulting mass, and afterwards adding nitric acid, which combines to the alkali, hydrous antimonic acid is also procured.

1924. Anhydrous antimonic acid is yellow, tasteless, and insolube water. When hydrous, it is white, and has the property of redden

litmus.

1925. Just at the moment when certain antimonies and antimonic subjected to a low red-heat, lose their water of crystallization, they gave to a transient light, as vivid as would result from a true combustion.

In the last edition of this work, I quoted, on the subject of tartar emulianticle previously published by Dr. Bache, in the American Cyclopedia of Prophedicine. This article I shall not introduce into this edition, because the infit tion which it comprises has been given in the United States Dispensatory, or we owe to my friend shovementioned, and to my colleague, Dr. Wood. I prophed that of this Dispensatory, every matriculant of our university will be provided a copy, as it appears to me to be of itself equivalent to a choice library of m medical knowledge.

they indur in cc no change in weight. Their colour is rendered brighter, and they become less susceptible of decomposition by acids. This result causes especially with the antimoniates of copper, cobalt, and zinc.

1926. The reaction of diluted nitric acid with antimony, is quite analogous to that already described in the case of bismuth. According to Berselius, a subnitrate results, which may be decomposed by water as already stated, and converted into a hydrated sesquioxide. But Thenard informs that, if this metal be subjected to nitric acid, it is converted into hydrous satimonious acid (acide antimonieux blanc et hydraté). Possibly the difference may arise from the acid being in one case concentrated, in the other lints.

Of the Compounds of Antimony with the Halogen Class.

1927. Sesquickloride of antimony may be obtained, as Thenard alleges, by distilling the metal with the bichloride of mercury; also by the reaction of aqua regia with metallic antimony, and subsequent distillation of the resulting liquid, collecting the product in a fresh receiver when it becomes oleaginous in its consistency. He recommends as preferable, however, the action of chlorohydric acid on the sesquisulphide with heat, allowing the sulphydric acid gas to escape into the fire. The resulting liquid is to be decanted, and concentrated by heat in a retort, until it acquires an aleaginous consistency.

1928. The sesquichloride has been designated as the butter of antimony. It is white, semitransparent, very caustic, fusible below a boiling heat, and rystallizable in tetrahedrons by refrigeration. It is volatile at a heat beow redness, and deliquescent, so as to be liquefied by exposure to air. It has already been mentioned, that by subjecting this chloride to copious affusions of water, (eight times its weight, according to Thenard) an oxy-

chloride results, formerly called the powder of Algaroth.

1929. Bickloride of antimony, agreeably to the last mentioned author,

exists only in combination with chlorohydric acid.

1930. Perchloride of antimony is formed by the combustion of the metal in chlorine. It is a yellow liquid, sending forth thick fumes into the air, with a strong and disagreeable smell. It attracts moisture, and is, in consequence, at first converted into a white crystalline mass, but afterwards liquefied by a further accession of humidity. Yet by exposure to a large quantity of water with heat, it is decomposed, and deposits hydrous antimonic acid. This process is recommended as the best for obtaining this compound.

Of the Compounds of Antimony with Sulphur and Selenium.

1931. It has been stated that antimony is procured principally from the sesquisulphide, which is found in the shops under the name of anti-

mony, the metal being distinguished as the regulus.

1932. Sesquisulphide of antimony may be formed from its ingredients, by heating the metal in a state of division with sulphur. It is more fusible than metallic antimony, is crystalline in texture, has a metallic lustre, and a bluish-gray colour. It may act either as a sulphacid, or as a sulphobase. With the sulphides of the alkalifiable metals it forms compounds which may be designated as hyposulphantimonites.

1933. The sesquisulphide and sesquioxide of antimony enter into combination with each other in different proportions, forming compounds which

must be called oxysulphides, consistently with the nomenclature adopted in

the case of the analogous compounds of oxides with chlorides.

1934. When the sesquisulphide of antimony is roasted, in other works exposed to heat with access of air, it becomes more or less exidized to cording to the duration of the exposure, the degree of heat, and the supply of air. If, after the roasting has continued for some time, the temperature be raised so as to fuse the mass, a vitreous compound will result, the composition of which will vary according to the ratio of the oxide to the subphide, at the time of effecting the fusion. According to Thornson, when the ratio of the former to the latter is as five to one, the compound has the name of crocus of antimony; when the ratio is as three to one, it has been called liver of antimony. This name, however, is given by Hernelson to a compound of the sulphides of antimony, with the sulphide of passaium or sulphide of sodium.

as it is melted, be kept for a great length of time in a state of fusion as carthen crucible, it derives a portion of exide of iron and silicic acid from the crucible, and thus forms a transparent mass of a yellow-hyacuth colour, commonly called the glass of antimony. This glass, according to Thenard, is a mixture of exysulphide of antimony, with the silicates of the silicates

timony and iron.

1936. By the reaction of the sesquisulphide of antimony with the allelies, either caustic or carbonated, and either in the wet or dry way, a material reaction ensues, by which the antimony of the sulphide is more or less sulphurized; while the residual sulphide of antimony, acting as a sulphacid, combines more or less with the resulting sulphobase of the alkalifiable metal.

1937. The extent to which the sesquisulphide, in the resulting sulphosalt, can be retained by the sulphobase in an aqueous solution, appears dependent upon temperature. Hence, whether the sulphosalt be produced to the dry way and dissolved in hot water, or be generated by horizing the agreements in this liquid, the sesquisulphide precipitates by refrigerature.

1938. The precipitate thus obtained, under the name of kerms was to was so much in vogue in France, about a century ago, as to induce the government of that country to purchase from a surgeon of the name of L.

Ligerie, the art of preparing it-

1939. Thenard alleges that it appears from the analysis of Henry, Inthat the composition of kermes varies according to the process employed for its production. When prepared by boiling the sesquisulphide in a shiftion of carbonate of potash or soda, kermes may be considered as a hydrated oxysulphide; but when procured by boiling the sesquisulphide in a solution of caustic potash or soda, or by fusion with them or their carbonates, and subsequent solution in hot water, it is a hydrated sesquisulphide containing very little if any oxide. As obtained by precipitation from that emetic by sulphydric acid, it is a pure hydrated sesquisulphide. After the kermes has precipitated, a portion of the sesquisulphide still remains a union with the sulphobase. Hence, on the addition of an acid, a further precipitation takes place, both of the sesquisulphide of antimony, and the sulphur of the sulphobase; and these, either by combination or analysis constitute the golden sulphur of antimony, another well known pharms constitute the golden sulphur of antimony, another well known pharms constitute the golden sulphur of antimony, another well known pharms contains a preparation.

1940. According to the analysis of Henry, Jr., as quoted by Thesart

omposition of kermes, when obtained in the wet way by carbonate of is as follows:

Seequisulphide of antimony,				•	•	62.5
Sesquioxi	de of a	ntimony,	•	•	•	27.4
Water,	•	•	•	•	•	10
Soda,	•	•	•	•	•	a trace.

41. Upon the whole it is inferred that the sesquisulphide, in precipity by refrigeration as abovementioned, combines with water in all cases; hat when the process is conducted in the wet way by means of a cared alkaline solution, the precipitating hydrated sesquisulphide combines the sesquioxide, forming an oxysulphide. The presence of carbonic in union with the alkali is requisite, in order to enable the menstruum m and dissolve while hot, a double carbonate of the alkali and sesquition. The latter, being thus taken up by the aid of heat, subsequently, precipitates in combination with this sulphide, as already men-

42. The officinal preparation, called precipitated sulphuret of antimony, ained by adding diluted sulphuric acid to a solution of the sesquisulof antimony in a hot solution of caustic potash. A precipitate rewhich may be considered as a mixture of kermes mineral and golden

ir of antimony.

ving antimonious acid in chlorohydric acid, and subjecting the resultquid to sulphydric acid. I infer that four atoms of chlorohydric acid, g on two atoms of antimony, in union with four atoms of oxygen, will aductive of a bichloride, and that this will be converted into a bisulphide action with the sulphydric acid.

44. The bisulphide, being resolvable into the sesquisulphide and sulby heat, cannot be produced by the fusion of its constituents. It is of

ange-red colour, and acts as a sulphacid.

5. Persulphide of antimony is obtained by passing sulphydric acid the addition of the perchloride of this metal, to which tartaric has previously been added. Its colour resembles that of the bisulphide, the somewhat paler.

46. The selenide of antimony is obtained by heating this metal with Like the sulphide, it is capable of entering into combination with

ade.

Experimental Illustrations.

47. Antimony and its sulphide, exhibited, and exposed e blowpipe: also, the crystals and solution of tartar ic. Kermes mineral, golden sulphur, and precipitated baret of antimony, exhibited. Antimony, subjected to a Kermes mineral, precipitated from a solution of are emetic by sulphydric acid.

SECTION XIII.

OF METALS PROPER OF MINOR IMPORTANCE.

OF PALLADIUM.

1948. Besides iron, copper, and lead, four metals, palladium, rhotom iridium, and osmium, are found in union with, or accompanying the sate grains of platinum, as imported from South America. Accordingly, is portion of that assemblage of metallic particles, of which the native grain of platinum above mentioned form the principal part, be digested a second regia, the platinum, together with the palladium, rhodium, copper, and will be dissolved; while a black powder will be left, consisting of committee.

and iridium in combination with each other.

by the chloride of ammonium, any palladium which it may contain, all of the other noble metals which may be present, may be precipitated in a bright plate of zinc. The resulting precipitate, after being digested to chlorohydric acid and washed with water, should be redissolved in aqua exception of acid should be neutralized by carbonate of sodu. From the neutralized solution the palladium may be thrown down by a solution of behanded of mercury which yields its eyanogen to the palladium. An unbedie cyanide of palladium, being thus formed, precipitates. By the additional this precipitate is decomposed, the cyanogen is expelled, and the metal is isolated.

1950. Mr. Cloud, of the United States' mint, found this metal in a name

alloy of gold which was brought from Brazil.

1951. The colour of palladium appears to me to have a minute degree of tendency towards the rosy hue of bismuth, not being quite so pale platinum, which it otherwise much resembles in appearance. It is however more fusible, rather harder, and more elastic. Its specific gravity, also, is much less, being about 11.5. It is malleable and ductile, and insusceptible of oxidizement by heat and air.

OF RHODIUM.

chloracids of rhodium, mercury, and several other metals, united with the chloride (or chlorobase) of sodium, resulting from the carbonate of size employed as abovementioned to neutralize the excess of acid. Those is likewise present a portion of the undecomposed bicyanide of integrity. Under these circumstances, chlorohydric acid must be added, a order to convert this bycyanide into a bichloride, and the solution are wards must be evaporated to dryness. The resulting mass should these washed with alcohol, which dissolves all the chlorosalts of sodium present except the chlorhodiate. Rhodium is obtained from this by heating it is current of hydrogen, which removes the chlorine combined with the mass the chloride of sodium being removed by water.

1953. Rhodium, according to Berzelius, cannot be fused, except by majecting it, when in the state of a sulphide or arsenuret, to an intense hat-After fusion, it resembles platinum in appearance. Its zalts are generally either red or yellow. It is named from its chloride, which is rose-red.

OF IRIDIUM.

954. When the black powder, consisting of the osmiuret of iridium, ch remains as above stated, after we have subjected the crude grains of inum to aqua regia, is heated with soda, an osmiate of soda is formed, ch may be removed by dissolving it in water. The remaining mass is e treated with aqua regia, in which the iridium, converted into a chlodissolves. By repeating this process, the whole is finally converted solutions of chloride of iridium, and of osmiate of soda.

955. From the former, crystals of the chloride of iridium may be obal by evaporation, which, on exposure to a strong heat, yield metallic

um.

956. Iridium resembles platinum in appearance, and is probably, acing to Thomson, the heaviest of the metals. When heated in conwith air nearly to redness it is oxidized, but on the application of a ser temperature it is again restored to the metallic state. Thenard, hower, states, that iridium which has been subjected to a strong heat, is absoly insusceptible of oxidizement by the air at any temperature.

957. Iridium is said to be the most refractory of the metals, having er been fused until it was placed between the poles of Children's large ranic battery. It was then converted into a globule, possessing metallic

tences and lustre.

OF OSMIUM.

958. Osmic acid may be obtained by distilling the solution of osmiate soda, procured as above described, with nitric acid at a gentle heat. comic acid passes over, and may afterwards be reduced by the additional passes over. It is, however, alloyed with mercury, mingled with the chloride of this metal. These may be sublimed by a

tle heat, leaving pure metallic osmium.

959. Osmium obtained in this way, is of a grayish-black colour; but portion of the volatilized oxide be made to pass with a current of hydrothrough a glass tube, the osmium is deposited in the form of a ring of allic brilliancy, and of a white colour. It is so difficult to fuse in close sels, and so liable to be volatilized when heated in the air, that it has y been obtained in powder, or in minute friable masses. Its volatility he air arises from its great susceptibility of oxidizement, and the volay of its oxide, the fumes of which are pungent.

OF NICKEL.

960. A mineral had been known to the German miners by the name supfer nickel, or false copper. About the middle of the last century, netedt alleged the existence, in this mineral, of a peculiar metal. Neveress, the metal, thus indicated, was considered by many chemists as an y of copper with iron. About 1775, Bergmann confirmed, by an iyais, the allegation of Cronstedt.

1961. Kupfer nickel is principally an arseniuret of nickel, but contains, salphur, iron, cobalt, and copper. Nickel is extricated from it by a

ious and intricate process.

1962. Nickel is of a white colour, difficult of fusion, malleable and not easiraidized by the air. It is so susceptible of the magnetic influence that a manent magnet may be made of it. If sufficiently abundant, nickel and be very valuable in the arts. A white alloy of this metal with copper, had long been known in China, under the name of packfong. Of his this alloy has been brought into use in Europe, under the name of argentane or German silver. It serves for pencil cases and many analogue uses nearly as well as silver. It combines with oxygen, chlorine, iodic, cyanogen, sulphur, and the metals. Its oxides are soluble in the acid, and in their habitudes are much like those of copper. The solubility of its protoxide in caustic ammonia, is an important means of separating nickel from its alloys.

OF CADMIUM.

1963. This metal has been derived only from the ores of zinc. During the reduction of calamine, a substance sublimes which yields from 12 to 20 per cent. of cadmium.

1964. A solution of the ore in sulphuric acid, being impregnated with sulphydric acid, the cadmium precipitates in the state of sulphide, mind with a little sulphide of zinc, and sometimes with sulphide of copper. When these sulphides are exposed to chlorohydric acid, the sulphur unites with the hydrogen of the acid and escapes, and they are converted into chlorida. Carbonate of ammonia being added to the resulting solution of cadmium and zinc, a carbonate of cadmium is alone precipitated. From this, in metal may be obtained by means of heat and charcoal.

1965. Cadmium is almost as white as tin, is without odour or take, very brilliant, and susceptible of a fine polish. It is crystallizable, malesble, and ductile, and so soft as to yield easily to a file or knife. Its specific gravity is 8.6 nearly. It is too scarce to be usefully applied. It fuses and

volatilizes at a very low temperature.

OF CHROMIUM.

1966. This metal is found in nature only in the state of an acid and of an oxide, generally united with lead or iron, though in some instances pure. It was in the native chromate of lead, found usually in crystals which rival the ruby in colour, that this metal was discovered by Vauquelin. A compound of the sesquioxides of chromium and iron, called incorrectly chromate of iron, is found plentifully in this country. The sesquioxide of chromium, when intensely heated with charcoal, is reduced, but not without great difficulty.

1967. The presence of chromium in a mineral may be detected by the fusion of a minute portion before the blowpipe with borax, or preferably, with the ammoniacal phosphate of soda. In this way, a globule of a beautiful emerald green results, which preserves its colour either in the oxidizage or reducing flame. By these characteristics it may be distinguished from copper or uranium; since uranium communicates a green hue only in the

reducing flame, copper only in the oxidizing flame.

1968. Chromium is a hard, brittle metal, of a grayish-white colour, and very difficult to fuse. Its specific gravity is 5.9. Its equivalent is 28. It forms with oxygen a sesquioxide and an acid. The compound, heretoice considered as a deutoxide, proves to be a mixture of sesquioxide and chromic acid.

1969. The sesquioxide of chromium is easily obtained by exposing the chromate of mercury to heat, by which the mercurial oxide and a portion of the oxygen of the acid are expelled, while the sesquioxide remains in the form of a grass-green powder. It may also be obtained in the state of hydrate, by mixing solutions of the bichromate of potash, and persulphing

ssium. This sesquioxide is of a beautiful green colour, which it nicates to some of its compounds, being in fact the colouring matter merald. It appears to act both as an acid and a base.

In common with zirconia and oxide of titanium, the sesquioxide mium, when obtained from the hydrate by expelling the water by a seat, becomes incandescent at a certain elevation of temperature, in which is altogether unaccountable. At the same time it loses its

y of solubility in acids which it before possessed.

chromic acid may be procured by the following process:—Let rts of the chromate of lead be mixed with three parts of fluoride of , both finely pulverized. Then let five parts of sulphuric acid, desor water as far as possible by boiling, be added, and let the whole led in a leaded or platinum alembic at a gentle heat. A red gas developed, producing in the air yellow fumes. This red gas is a of chromium, which, on being passed into water, is converted into ric and chromic acids. The former may be expelled by evaporate chromic acid remaining in a state of purity.

If, instead of causing the gaseous fluoride of chromium to enter it be conducted by means of a tube into a receptacle of platinum, with moistened paper, and having a small quantity of water at the the gas will be decomposed by the aqueous vapour, mingled with of the vessel, and will deposite first about the mouth of the tube, and rds throughout the vessel, a flocky vegetation, consisting of ruby-

stals of chromic acid.

. Chromic acid is solid, soluble in water, and capable of reddening It is decomposed by heat, and by most substances which possess nity for oxygen. It possesses an acid and astringent taste, and a d colour, which it communicates to some of its compounds, as aloticed in the case of native chromate of lead. It forms striking and I precipitates with various metals. That which it produces with of a splendid orange-yellow, and is much used as a pigment. of the streak left by the red crystals above described, when rubbed hard surface, is likewise orange-yellow; and the same change enm pulverization. The bichromate of potash is poisonous, and no se acid and its compounds are generally poisons. Chromic acid a stain upon the skin which cannot be removed by water, unless it an alkali. Where there is any abrasion of the cuticle, the presence acid will induce a painful ulcer. Hence the sores to which dyers osed who employ bichromate of potash as a dye-stuff. These sores en alleged to arise even from exposure to the vapours or fumes of d. When received into the stomach, chromic acid is a virulent

Dr. Ducatel informs us of the case of a labourer who died in five after drawing into his mouth from a syphon, a solution of bichromate sh; although he was under the impression that, by spitting, he had

taking it into his stomach.

Dr. Ducatel suggests an alkaline solution as the best antidote for this us salt; as he ascribes its activity mainly to the excess of acid. An e of a criminal prosecution for poisoning by this bichromate, is menwhich failed from that ignorance of its deleterious properties which estel's communication must tend to correct.*

Journal of the Philadelphia College of Pharmacy, for January, 1834, page 5.

OF COBALT.

1975. This metal is found in nature, principally in union with arms. By the exposure of the mineral, thus containing it, to heat, with access of air, the arsenic is exidized and expelled, and the cobalt is reduced to the state of an impure exide, called zaffre. By fusion with the alkah and sand, zaffre yields a beautiful blue glass, which, when pulverized, form the blue vitreous powder called smalt.

1976. Cobalt may be obtained from its oxide, by intense ignore with charcoal, or by subjecting it, while ignited in a porcelam tube, to a outsile

of hydrogen.

1977. Cobalt is brittle, of a grayish-white colour, and feeble lustra be specific gravity is 8.5 nearly. It requires a high temperature for its functions.

OF COLUMBIUM.

1978. A metal discovered by Hatchett, in 1801, in a mineral obtains from America, received the name of columbium. It was afterwards detected by Ekeberg in two Swedish minerals, called tantalite and yttrotantames, and being supposed to be a new metal, was called tantalum. Wollasto afterwards demonstrated the identity of tantalum with columbium.

1979. This metal is found in the state of an acid, combined rither manganese and a little iron, or with yttria. Both combinations are way rare. It may be reduced by heating the fluocolumbate of potassius.

fluoride of columbium and potassium, with potassium.

1980. Columbium is a brittle metal, of an iron-gray colour, having the metallic lustre. It is infusible by the most intense heat of the forge fire-

OF MANGANESE.

1981. Manganese exists in nature principally in the state of a black bioxide; rarely in that of phosphate, sometimes in the state of sulphot. The utility of this oxide, as a source of oxygen gas, as an ingredient in glass, and as one of the agents for evolving chlorine, has been noticed. (663.) The metal is obtained by heating the oxide intensely with charmal or potassium. It is gray, brittle, hard, and scarcely fusible by the highest heat of the forge, or air furnace. In the metallic state, it has not been applied to any useful purpose.

1982. Manganese is remarkable for the number of compounds which a forms with oxygen. Besides a protoxide, sesquioxide, and bioxide, it forms two acids, the manganic, and oxymanganic or permanganic acids.

salts of the latter detonate with combustibles.

1983. When the black oxide (bloxide) is fused with nitrate of potah, a compound results, of which the aqueous solution becomes blue, violet, and red, and finally colourless. Hence this compound has been called charaltern mineral. These colours appear to be produced by the conversion of the manganate of potash, into an oxymanganate.

OF MOLYBDENUM.

1984. This motal is only found in the state of sulphide, resembling planbage, or united with exygen and lead in the state of molybdate of leadfrom the sulphide it is obtained by ebullation with nitric acid, which acidise both the sulphir and metal. The sulphiric acid being expelled by him the molybdic acid is decomposed by intense ignition with charcoal. 985. As from the difficulty of fusing it, molybdenum has been only ined in small grains, its properties are but little known. It is alleged ave a high degree of metallic lustre, and a white colour.

OF TITANIUM.

e knowledge which human skill and assiduity have accumulated, with ct to the materials of the globe which we inhabit. It is obtained by rating the oxide from the substances with which it is naturally mixed, neating it intensely with charcoal.

87. Titanium was first ascertained to exist in the state of oxide, by 3regor, in a mineral called menachanite. It was subsequently detected e metallic state by Dr. Wollaston, in minute cubic crystals, in the slag

d at the bottom of a smelting furnace.

188. These crystals were conductors of electricity, of the specific graof 5.3, and hard enough to scratch rock crystal. In colour and lustre, were like burnished copper. They resisted the action of nitric acid aqua regia, but were oxidized by being heated with nitre.

OF TUNGSTEN.

169. In 1781, Scheele, having analysed a stone known by the name ngsten, or heavy stone, concluded that it consisted of an acid united lime. Bergmann suspected the radical of this acid to be metallic. Its. D'Elhuyart verified his conjecture, by heating tungstic acid intensely charcoal.

390. Transsten is grayish-white, brilliant, and extremely difficult to Its specific gravity is 17.6.

OF URANIUM.

191. Uranium is a rare production in nature, and has scarcely been in sufficient quantities for an adequate observation of its properties. stated to have the metallic lustre, a reddish-brown colour, to be cryste in its structure, and scarcely susceptible of fusion by the heat of a fire.

OF CERIUM.

192. Cerium, according to Vauquelin, who was unable to obtain it in the larger than the head of a common pin, is a white brittle metal. I some experiments made by Children and Thomson, it appears to be sptible of volatilization.

OF VANADIUM.

203. Vanadium was discovered, in 1801, by Del Rio, in a lead ore Zimapan, in Mexico; but Collet Descotils, to whom the mineral was having made some new experiments upon it, pronounced it to be an of chromium. Del Rio himself having acquiesced in this opinion, it generally adopted until 1830, when Sefstrom discovered this metal n in a variety of Swedish iron, and in the scoria of the forge at which iron had been wrought.

194. Vanadium resembles molybdenum in appearance; and in its pro-

as lies between that metal and chromium.

Experimental Illustrations.

1995. Exhibition of various specimens of the metals mentioned in the preceding pages. Magnetic influence of nickel, demonstrated. Solutions of silver, mercury, and lead, precipitated by chromate of potash. Sesquionide of chromium, evolved by heating the chromate of mercury. Exhibition of the fluoride of chromium. Effects of co-balt, also of manganese, upon vitrified borax.

SALTS.

and nomenclature adopted in this work, I alleged the word salt to be more ceptible of any definition consistent with the use made of it by Bernelius a basis of nomenclature. As the reader, who has studied this work so fit, as to have reached this page in due course, should have acquired a knowledge of the facts upon which the above cited allegation was founded, I will here quote the language in which those facts were stated, and my inference.

from them justified.

1997. The most striking feature in the nomenclature of Berzelius, is the formation of two classes of bodies; one class called "halogene," or sik producing, because they are conceived to produce salts directly; the other called "amphigene," or both producing, being productive both of acids and bases, and of course indirectly of salts. To render this division charle, it appears to me that the terms acid, base, and salt, should, in the first place, be strictly defined. Unfortunately there are no terms in use, more broad, vague, and unsettled in their meaning. Agreeably to the common acceptation, chloride of sodium is pre-eminently entitled to be called a salt : same in common parlance, when no distinguishing term is annexed, tak is 🌬 name of that chloride. This is quite reasonable, as it is well known that the genus was named after this compound. Other substances, having their obvious qualities some analogy with chloride of sodium, were, at an early period, readily admitted to be species of the same genus; as, for a stance, Glauber's salt, Epsom salt, sal ammoniac. Yet founding their protensions upon similatude in obvious qualities, few of the substances called salts, in the broader sense of the name, could have been admitted into the class. Insoluble chlorides have evidently, on the score of properties. little claim to be considered as salts, as insoluble exides. plumbum corneum, butter of antimony, and the furning liquor of Laburan, are the appellations given respectively to chlorides of silver, lend, antimosy, and tin, which are quite as deficient of the saline character as the comsponding compounds of the same metal with oxygen. Fluoride of calculations (fluor spar) is as unlike a salt as lime, the oxide of the same metal. He saline quality can be perceived in the soluble "halord salts," so called by Berzelius, while free from water; and when a compound of this kind is

BALTS. 357

e, it may be considered as a salt ned of an hydracid and an oxyt of the water with the halogene and of the oxygen with the cal. It is admitted by Berzelius, and of the oxygen with the cals. It is admitted by Berzelius, and of the oxygen with the constrated that the elements of to the constrated that the elements of to the constrated that the elements of to the considered as a salt of the hydrocolous of the hydrocolous of the hydrocolous of the hydrocolous of the oxygen with the constrated that the elements of to the considered as a salt of the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of to the oxygen with the constrated that the elements of the oxygen with the constrated that the elements of the oxygen with the constrated that the elements of the oxygen with the constrated that the elements of the oxygen with the constrated that the elements of the oxygen with the constrated that the elements of the oxygen with the constrated that the elements of the oxygen with the constrated that the elements of the oxygen with the constrated that the

as the criterion of a salt; if, as in some of the most respectable chell treatises, we assume that the word salt is to be employed only to deate compounds consisting of a base united with an acid, we exclude the class chloride of sodium, and all other "haloid salts," and thus set the basis of the distinction between "halogene" and "amphigene" sents.

999. Moreover, while thus excluding from the class of salts, substances ch the mass of mankind will still consider as belonging to it, we assemunder one name combinations opposite in their properties, and destitute he qualities usually deemed indispensable to the class. Thus under the nition that every compound of an acid and a base, is a salt, we must ch this name to marble, gypsum, felspar, glass, and porcelain, in comwith Epsom salt, Glauber's salt, vitriolated tartar, pearlash, &c. But itting that these objections are not sufficient to demonstrate the absurdity isfining a salt, as a compound of an acid and a base, of what use could a definition be, when, as I have premised, it is quite uncertain what is scid, or what is a base. To the word acid, different meanings have a attached at different periods. The original characteristic sourness is longer deemed essential! Nor is the effect upon vegetable colours treated un indispensable characteristic. And as respects obvious properties, can re be a greater discordancy, than that which exists between sulphuric 1, and rock crystal; between vinegar, and tannin; or between the vola-, odoriferous, liquid poison, which we call prussic acid, and the inodos, inert, concrete material for candles, called margaric acid?

1900. While an acid is defined to be a compound capable of forming a with a base, a base is defined to be a compound that will form a salt han acid. Yet a salt is to be recognised as such, by being a compound the acid and base, to which, as I have stated, it is made an essential

an of recognition.

3001. An attempt to reconcile the definitions of acidity given by Berze, with the sense in which he uses the word acid, will, in my apprehen-

1, increase the perplexity.

1002. It is alleged in his Traité, page 1, Vol. II., "that the name of it is given to silica, and other feeble acids, because they are susceptible combining with the oxides of the electropositive metals, that is to say, it salifiable bases, and thus to produce salts, which is precisely the neighborhood character of acids." Again, Vol. I., page 308, speaking of the logene elements, he declares that "their combinations with hydrogen, are tony acids, but belong to a series the most puissant that we can employ chemistry; and in this respect they rank as equals with the strongest of acids, into which oxygen enters as a constituent principle." And again, I. II., page 162, when treating of hydracids formed with the halogene sa, he alleges "The former are very powerful acids, truly acids, and feetly like the oxacids; but they do not combine with salifiable bases; the contrary, they decompose them, and produce haloid salts."

2009. In this paragraph, the acids in question are represented as preeminently endowed with the attributes of acidity, while at the same they are alleged to be destitute of his "principal character of acids," the

property of combining with salifiable bases.

volumes of oxygen and one of nitrogen, considered by chemists generally as a distinct acid, Berzelius uses the following language. "If I have not coincided in their view, it is because, judging by what we know at present the acid in question cannot combine with any base, either directly or university; that consequently it does not give salts, and that salifiable have decompose it always into nitrous acid, and nitric oxide gas. It is not then distinct acid, and as such, ought not to be admitted in the nomencial viewing these passages with all that deference which I feel for the productions of the author, I am unable to understand upon what principle the colusion of nitrous acid from the class of acids, can be rendered constant with the retention, in that class, of the compounds formed by hydrogen

with " halogene" elements.

2005. It is certainly to be regretted that there should be so much different culty in giving a precise meaning to a word used so extensively as the which led to the language above quoted. The best definition which I a devise, in this case, is that a salt is a compound, resulting from the una of at least two acid, acrid, or corrosive ingredients; forming, agreeably the language of the older chemists, a tertium quid, or in plain English. third something, differing materially from its constituents. It should as I conceive, be crystallizable, and soluble either in water or alcohol. 1 de not think that a satisfactory line of demarcation can be drawn between salts, acids, and bases. Some compounds which lean so much toward salinity in their characteristics, as to have been classed with salts, have latterly been found to play the part of acids or bases, as instanced by the binary halogen salts. I would consider them as salts when acong as such and as acids or bases when acting as acids or bases. Bermins has soggested this kind of contingent definition in the instance of water: which he represents as acting as a base with some acids, and as an acid with some bases. Thus it seems possible for the same body to act cither as an acce. a salt, or a base, accordingly as it may be associated.

Of the Principal Groups of Salts.

2006. As respects composition, I conceive that there are at least three groups of salts.

2007. 1st. Binary saline compounds of a halogen ele-

ment and a metal.

2008. 2d. Saline compounds of acids and bases, tertion quids agreeably to the definition of acidity and basidity. (631.)

2009. 3d. Saline compounds containing either an organic acid or an organic base; or consisting of such an

acid, united with such a base.

[&]quot; I am unable to refer to any authority for the use of this word, but concern myself justified in employing it, as, by analogy, it cannot be misunderstood by the reader.

). As far as consistent with the due allotment of I have given an account of the first group in treatthe metals. But the class thus constituted are e of combining with each other, and with the electrore or acid compounds, formed by the union of their n ingredients with non-metallic radicals. In this empounds are produced, which Berzelius designates ble haloid salts. I, however, consider them as much l to be treated as saline compounds of acids and as the double sulphides, selenides, or tellurides, are so treated by that distinguished chemist. (627.) 1. I shall designate the salts comprised in the 1st groups abovementioned by their basacigen ingredi-633.) Hence the nine following classes; oxysalts, alts, selenisalts, tellurisalts, chlorosalts, bromosalts, is, fluosalts, and cyanosalts. Obviously, of these the ir are formed by the amphigen bodies, and the rest halogen bodies of Berzelius.

SECTION L

OF OXYSALTS.

2. In describing the oxysalts, I shall be constrained fine my remarks to some of the more important teristics of each of the sets of salts formed by the it inorganic oxacids with the more energetic bases. If those formed by acids of minor importance will tted altogether.

Of Chlorates and Hypochlorites.

3. As agreeably to the premised arrangement, the 3, first made the objects of attention in this work, nose formed by the union of chlorine with oxygen; was that the saline combinations formed by those with bases, should be the first to be treated of oxysalts.

decies of salts, hypochlorites, chlorites, chlorates, exchlorates, or oxychlorates. It seems, however, whether chlorous acid can be presented to a base, t being resolved into a chlorate and chlorine. In spect, it seems to rest on the same footing as ni-

trous acid. (984, 2004.) Of course there are no diorites.

2015. The chlorates and hypochlorates are the product of one process, in which an oxybase is made to react whether in the chlorine. In the process alluded to, a fixed alkali, or the of the three more powerful alkaline earths, whether in the pulverulent state of a hydrate, being sufficiently exposed to chlorine, is for to acquire the bleaching and disinfecting properties which that gas is so remarkably endowed.

2016. In the case in which a solution of potasses saturated with the gas, besides the acquisition of bleach power, by the mother water, crystals result of chlorated potash, which from their inferior solubility precipitate.

ment, when first employed; because, agreeably to the science of the day, nothing could have been less likely succeed. At that time, chlorine was considered as a oxacid of an unknown radical. (886.) But if the bleading and disinfecting properties of chlorine were due to acidity, nothing could be less consistent with the retention of those properties, than saturation with powerful base Subsequently, when the elementary character of chlorine became known, the ascertained retention of its bleaching and disinfecting power, after combination with an orphase, appeared much more consistent with the suppose nature of the ingredients.

2018. It was conceived, that chlorine feebly attracte to an oxybase, was liberated by its affinity for colours matter, or feculent emanations, or by the affinity of an other principle for the oxybase. Accordingly, until with the last ten years, the impression generally prevailed, to the liquids, powders, or salts employed in bleaching, we compounds of an oxybase with chlorine. Hence, the terms chloride of lime, or chloride of potassa, or of some which are still in use, especially among manufacture chemists.

2019. It was in the treatise of Berzelius, that I first with the explanation which I gave in the last edition this text book, of the process under consideration, which I now subjoin.

2020. When into a solution of potash, (oxide of potasium,) chlorine is introduced, one portion of it combined

with the potassium, separating from each atom the atom of oxygen by which it was oxidized. The oxygen thus liberated from several atoms of the metal, coming into contact with another portion of chlorine, forms with it thlorous acid. Each atom of the acid, thus formed, unites with an atom of potash, producing a chlorite. By con-inuing the operation until all the potash which remains bee is decomposed, that which has meanwhile united with he acid is attacked by the chlorine, and the oxygen, libeated in consequence from each atom of the chlorite, conrerts another portion of this salt into a chlorate. n crystals, which being subjected to boiling water, are purified by the recrystallization which cooling induces.

2021. This explanation seems to require modification, only so far as to introduce the hypochlorous in lieu of the chlorous acid, (688,) agreeably to the new view of the subject presented in treating of the compounds of chlorine with oxygen. Reference is there made to the investigaions of Balard, by which it has been shown that the gaseous product supposed to be the protoxide of chlorine, called euchlorine by Davy, is really a mixture of chlorous acid with chlorine; and also that the real protoxide of chlorine, is the acid which is formed during the process for making chlorate of potash, or bleaching powders, and which is now designated as hypochlorous acid.

2022. The impure hypochlorite of lime, called bleaching salt, is obtained by exposing hydrate of lime to chlorine. Analogous salts of potash and soda are found in the mother waters of the chlorates of those alkalies, and may likewise be obtained, by double decomposition, from the hypochlorite of lime, and carbonate of potash or soda. When obtained by these methods, hypochlorites are mingled with the chlorides of the metals peculiar to their

respective bases.

2023. Properties.—The hypochlorites are extremely susceptible of decomposition. This, however, takes place in a manner which varies with the circumstances in which they are placed. Bleaching or disinfection is effected by them when quite pure, by imparting oxygen; being resolved into this element and a chloride. Chlorine produces similar effects, by causing water to impart oxygen. No doubt the result is the consequence of complex affinity,

the hydrogen being attracted by the chlorine, the oxygen

by some oxidizable substance.

2024. When carbonic acid has access to an hypochorite, it combines with the base of the salt, displacing the acid; and if a chloride be present, its radical is oxidized by the oxygen of the acid thus displaced; while its chlorine is liberated, as well as that of the chlorous acid. Of course an evolution of chlorine must ensue from the employment, in like case, of any acid, which, in its uffinities, is not less energetic than carbonic acid. If, however, a pure hypochlorite, formed by the action of hypochlorous acid on a base, be subjected to the action of a more powerful acid, the hypochlorous acid may be liberated without being decomposed.

2025. When an aqueous solution of a hypochlorite is boiled in pure water, one portion of the chlorite is coverted into a chloride; while the oxygen, which is liberated from it during this transformation, converts another

portion into a chlorate.

2026. According to Thomson, when chloride of ammonium is introduced into a retort containing the hypochlorite of lime of commerce, made into a paste with water, gaseous nitrogen is evolved with a reaction so violent, that, in order to delay the extrication until his arrangements for collecting the product were completed, he found it expedient to wrap the hypochlorite in blotting paper.

Experimental Illustrations.

2027. Production of hypochlorite of lime. Its effects upon colouring matter. Evolution of nitrogen from chloride of ammonium, by hypochlorite of lime.

Properties of the Chlorate.

2028. The chlorates resemble the nitrates in deflagrating with combustibles; but the residuum which they leave is always a chloride; and the deflagration is more powerful and more easily effected. If chlorate of potash be triturated in contact with sulphur or phosphorus, an explosion ensues. Salts of this class give up their oxyges, and are converted into chlorides, simply by being heated. They are almost all soluble in water. The chlorate of the protoxide of mercury is said to be but sparingly soluble.

2029. The properties of the chlorates are most conniently illustrated by the chlorate of potash, which is an ficient material in several varieties of the matches which e ignited either by contact with sulphuric acid, by fricon, or crushing.

2030. Alcohol, or any of the essential oils, oil of turpenie for instance, may be ignited by means of chlorate of

tash and sulphuric acid.

Experimental Illustrations.

2031. Ignition of phosphorus with chlorate of potash percussion. Explosion of sulphur mixed with the chlote, by trituration. Composition for friction matches consting of chlorate of potash, sulphur, and phosphorus, ingled with gum, exhibited and ignited. About as much lorate of potash as may be piled upon a half cent, being posited in a heap amid the inflammable liquid, the afsion of concentrated sulphuric acid upon the heap, causes e liquid to be inflamed.



Combustion of Phosphorus under Water, by means of Chlorate of Potash and Sulphuric Acid.

2032 Let there be two tubes, one within the other, as here represented; the larger one, closed at the lower end, and containing water; the other open at both ends, the upper orifice funnel-shaped, and the bore about one-twelfth of an inch in diameter. Allow some very small pieces of phosphorus, and a few crystals of chlorate of potash, to fall down to the bottom of the large tube. Then, into the smaller tube, pour some sulphuric acid; so that, without being much diluted, it may descend upon the chlorate and phosphorus. A vivid ignition ensues, in despite of the surrounding water

2033. The sulphuric acid, uniting with the potash of the chlorate, liberates chloring and oxygen, which, coming into contact with the phosphorus, cause its

coming into contact with the phosphorus, cause its

combustion

Of Perchlorates or Oxychlorates.

2034. One of the processes for procuring oxychlorate of potash, has been mentioned in the text, while treating of oxychloric acid, (712,) and another is mentioned in a note Oxychlorates of other bases, are obtained either by double decomposition; or by the direct union of the acid, made as already suggested. (713.)

2035. The oxychlorates, in general properties, resemble the chlorates. They appear, however, to be less susceptible of decomposition; since the oxychlorate of potash is not decomposed by any of the acids at ordinary temperatures, and does not react as violently with sulphur as the

chlorate of potash.

2036. Nearly all of the oxychlorates would appear to be deliquescent, and soluble in alcohol, excepting those of potash, lead, protoxide of mercury, and ammonia. At the temperature of 59°, oxychlorate of potash requires for its solution sixty-five times its weight of water.

OF NITRATES.

2037. This class of salts is distinguished by deflagrating with charcoal and other combustibles. When the combustible is susceptible of acidification, the resulting acid unites always with the base. Thus in the case of charcoal, a carbonate is left; in the case of silicon, a silicate; in the case of sulphur, a sulphate; in the case of arsenic, an arseniate. They differ from the oxysalts containing an acid with a halogen radical (the chlorates for instance,) in leaving an oxide after deflagration, instead of a haloid salt. Thus the nitrate of potash leaves the oxide of potassium; while the chlorate leaves a chloride of potassium.

2038. If subjected to concentrated sulphuric acid, the nitrates, when dry, emit fumes of nitric acid. When added to liquid chlorohydric acid, by dehydrogenating the chlorine, they enable it to act on gold leaf, as it does when

presented to this metal in aqua regia-

2039. The neutral nitrates are all soluble, and many of them deliquescent.

Experimental Illustrations.

2040. Deflagration of a nitrate upon ignited charcoal and of charcoal and other substances upon fused nitrate of

otash, soda, copper, or strontia. A nitrate added to quid chlorohydric acid containing gold leaf, causes the olution of the metal. Decomposition of a nitrate by eat.

OF NITRITES AND HYPONITRITES.

2041. It would appear that the compound, which, conistently with the practice of the British and French chemsts, has been designated as nitrous acid, is decomposed then presented to bases, forming a nitrate and hyponitrite. t is probable, therefore, that there are no salts which roperly deserve the name of nitrites, in the sense in which his appellation has been used by the chemists abovemenioned. It has already been stated that Berzelius does not idmit the existence of any acid intermediate, as respects he quantity of oxygen contained, between nitric and hyponitrous acid, and, therefore, calls the acid to which the ast mentioned name has been applied, nitrous acid, and of course designates its compounds as nitrites. (984.) The hyponitrites of the English and French chemists, or utrites of Berzelius, resemble the nitrates in most of their properties; but may be recognised by the red vapours which they evolve on the addition of any of the stronger icids. (981, &c.)

OF SULPHATES.

2042. Their solutions all yield precipitates with solutions of baryta. Heated in contact with charcoal or hydrogen, they are converted into sulphurets, which, if moistened, smell like rotten eggs. They are almost all insoluble in alcohol. The sulphates of baryta, tin, antimony, bismuth, and lead, are quite insoluble in water. Those of strontia, lime, yttria, mercury, silver, and the sesquioxide of cerium, are nearly insoluble; while all other sulphates are soluble.

Experimental Illustrations.

2043. Precipitation of sulphates by solutions of baryta. Conversion of a sulphate into a sulphuret before the blow-ipe, demonstrated by the subsequent effect upon a metal-c solution.

OF HYPOSULPHATES, SULPHITES, AND HYPOSULPHITES

2044. The hyposulphate of baryta, is obtained by adding sulphate of barium to a solution of hyposulphate of manganese. (764.) The hyposulphates of lime and strontia are produced in the same manner, and to hyposulphates of the other bases, either by double decomposition, or by adding the acid directly to the base.

2045. The neutral hyposulphates are probably all soluble. The property, together with their conversion into sulphates by heat, and the court of sulphurous acid which they emit on the addition of concentrated sulphurous acid which the addition acid which the addition acid which the addition acid which the addition aci

phuric acid, is sufficient to enable us to recognise them-

2046. The insoluble sulphites are obtained by double decomposites; those which are soluble, by the direct action of the acid on the base.

2047. The sulphites are generally insoluble, and may be recognised by the odour of sulphurous acid which they coult on the addition of disal sulphuric acid; while they do not, like the hyposulphites, simultaneous deposite sulphur: also by their not yielding, like the hyposulphates, a total sulphate by heat.

2048. The hyposulphites are procured by treating metallic zinc, iron, a manganese, with liquid sulphurous acid. Each atom of this acid abnormal one atom of oxygen to the metal, being consequently converted into hyposulphurous acid, which, with the resulting oxide, forms a hyposulphure.

2049. The hyposulphites may likewise be formed by boiling suplies with flowers of sulphur, by which each atom of acid in any sulphite uto up an additional atom of sulphur, converting the sulphite into a hypostephite.

2050. The hyposulphites may all be decomposed by best, and, what acted on by sulphuric acid, deposite sulphur and liberate sulphusous acid.

OF SELENLATES.

2051. The seleniates greatly resemble the sulphates in properties. They are in fact isomorphous with them, and crystallize with the same quantity of water of crystallization. The seleniates are, however, more susceptible of decomposition than the sulphates, and when thrown on burning could deflagrate.

OF PHOSPHITES.

2052. The phosphites are obtained either by presenting the acid directly to the base, or by double decomposition. When thrown on burning coals they produce a yellow flame, the colour of which increases in interesty with the quantity of acid contained in the salt.

OF PHOSPHATES.

2053. The phosphates all give precipitates with solutions

of baryta, lime, lead, and silver.

2054. The phosphates are not decomposable by host alone. Those of the metals proper may be converted, by heat and charcoal, into phosphurets of the metals pecular to their respective bases. In the case of the phosphate of the earths and alkalies, a portion of the phosphoric and

deoxidized by the carbon, evolving phosphorus; while e remainder forms with the base a subphosphate.

2055. By heat the phosphates are converted into paraosphates, identical in composition, though different in

operties.

2056. According to Thenard, phosphoric acid combines th bases in five different proportions, forming biphosates, sesquiphosphates, neutral phosphates, sesquibasic phosates, and bibasic phosphates, in which the equivalents of id to those of the base are respectively as 2 to 1, 1½ to 1, to 1, 1 to 1½, and 1 to 2.

OF CARBONATES.

2057. This class of salts is distinguished by being susptible of decomposition, with effervescence, by any of e acids, excepting a few that are remarkably feeble, as, r instance, the cyanhydric and meconic acids.

2058. All the alkaline carbonates are decomposable by at, excepting those of potassa, soda, baryta, strontia, and

obably lithia.

2059. Each of the alkalies, potash, soda, and ammonia, rms with carbonic acid, a carbonate, consisting of an juivalent proportion of each ingredient; a sesquicarbone, in which there are one equivalent and a half of acid one of alkali; and a bicarbonate, in which there are of equivalents of the acid to one of alkali. When satuted with the acid, they are more susceptible of crystalization, and less nauseous to the taste.

2060. The evolution of the acid from the carbonates of

ne and ammonia has been already exhibited.

OF BORATES.

2061. The biborate of soda is found in nature in certain lakes, and is own in commerce as borax. In the examination of minerals by the

wpipe, it is very useful.

2062. The other soluble borates, which are those of potash, soda, lithia, d ammonia, are obtained by uniting the acid directly with the base. The rates, which are quite or nearly insoluble, are procured by double demposition with the borate of soda. Borates are undecomposable by heat, ben their bases are undecomposable by that agent. Other borates, when tensely heated, are resolved into oxygen, a metallic radical, and boric id.

OF SILICATES.

2063. The silicates are procured either by double decomposition, or by thing silicic acid strongly with the base. They are not decomposable by

heat alone; although, when heated with charcoal, some of the silicates are converted into silicurets. All the silicates, excepting those of potash, sale, and lithia, are insoluble.

OF CYANATES AND FULMINATES.

2064. The soluble cyanates are decomposable by water, and, if insoluble, by acids, into carbonic acid and ammonia. The fulminates are chiefly remarkable for the violent explosions which they produce by heat or percussion. The fulminate of mercury is employed as priming in percussion gun locks. It may be obtained by the following process: Dissolve 100 grains of mercury with heat in a measured ounce and a half of nitric acid of moderate strength; when cold, mix the solution with a measured ounce and a half of alcohol, and apply heat till effervescence takes place. When red fumes appear, check the action with water. The powder which peripitates, well washed with water, and afterwards dried at a gentle heat, will be the fulminate of mercury.

OF DOUBLE OXYSALTS.

2065. There are many cases in which two salts, formed by different bases but of the same acid, enter into combination. A compound thus constituted, formerly received the appellation of a triple salt, but is now designated as a double salt.

2066. Tartar emetic is a double tartrate, consisting of tartrate of potential

combined with tartrate of antimony. (1919.)

2067. Rochelle salt is a compound of tartrate of potash with the tartrate of soda. An analogous compound is formed by the union of tartrate of potash with tartrate of iron, called ferri et potassæ tartras, or tartrate of potash and iron, in the United States' Dispensatory; to which I refer stadents for much valuable information which my limits will not allow me to add.

2068. The saline compound, well known under the name of alum, is composed of one atom of trisulphate of alumina, and one of sulphate of pot-

ash, besides twenty-four atoms of water of crystallization.

2069. Other double sulphates have been formed analogous to alum, substituting soda or ammonia for potash, or iron, manganese, or chromium in alumina.

2070. Double silicates and carbonates exist in nature. Dolomite is a species of marble, consisting of the carbonates of lime and magnesia in equivalent proportions. Felspar consists of a silicate of alumina and a silicate of potash. Many native crystals, well known to mineralogists, are double silicates.

2071. Glass, in general, is a combination of one or more silicates. Figures, according to Turner, is a double sexsilicate of lead and potash.

2072. It ought not to be supposed that double salts are always projectly by the combination of single salts previously existing separately. In the case of tartar emetic, the bitartrate of potash, containing two equivalents of acid to one of base, is converted into the double tartrate of potash and and mony, by saturating with one equivalent of the sesquioxide of this means one equivalent of the acid in the bitartrate. Thus a tartrate of antimony produced in combination with a tartrate of potash, and a double salt is a course formed.

2073. It appears possible for two double salts to combine, as when the

soda (borax) is added to bitartrate of potash, in order to produce soluble cream of tartar" of pharmacy. According to Berzelius, this und may be considered as consisting of a double tartrate of potash da (sal Rochelle), combined with a double tartrate of potash, and cid acting as a base. See United States' Dispensatory.

SECTION II.

OF SULPHOSALTS.

1. Berzelius alleges that the metallic sulphides, which are capable bining with each other to form sulphosalts, contain for each atom of , the same number of atoms of sulphur, as the salifiable oxybases tacids of the same radicals contain of oxygen. In consequence of alogy in composition, if sulphydric acid gas be transmitted through entrated solution of an oxysalt, in which the acid and base have metallic radical, the hydrogen of the sulphydric acid takes all the 1 from both radicals. Meanwhile, an equivalent number of atoms hur, consequently liberated, take the place of the oxygen, forming a salt, consisting of a sulphacid and a sulphobase, analogous, in the r of atoms of each ingredient, to the oxysalt, from the decomposiwhich it originates.

5. In order, however, to effect the combination of the electro-positive c sulphides which act as bases, with the sulphides of non-metallic s which act as sulphacids, a different method must be adopted. e of sulphydric acid gas, which does not combine, except with the les of the metals of the alkaline earths and alkalies, it is either it into contact with a carbonate of the base heated to redness, or ade to enter into a solution of the hydrate. Whichever method be I, no access of atmospheric oxygen should be allowed. In either ne portion of the sulphydric acid is decomposed, its hydrogen comwith the oxygen of the base, and its sulphur with the metal; while per portion of the acid unites with the sulphide thus formed, proa sulphydrate.

8. It has been stated, (1248,) that combinations of sulphocarbonic may be formed with most of the electro-positive sulphides, either by mion, or by double decomposition. There are other methods of prethese sulphosalts, of which I cannot treat, consistently with the

prescribed for this work.

SECTION III.

OF SELENISALTS AND TELLURISALTS.

7. As has been already stated, both selenium and tellurium are e of combining with different radicals, forming selenides and tellu-These, in many cases, like the corresponding compounds formed by r, unite together to form selenisalts and tellurisalts. The resulting ations, however, have been but little studied.

SECTION IV.

OF CHLOROSALTS, BROMOSALTS, IODOSALTS, AND FLUOSALTS OF THE SECOND GROUP.

2078. The chlorosalts are generally obtained by mingling chloracit with chlorobases (631), either in the wet or dry way. In the latter case, heat must be employed in order to facilitate their union.

2079. The bromosalts and iodosalts may in general be obtained a the same manner, by mingling bromacids with bromobases, or iodacals with

iodobases.

2080. I have mentioned, in treating of the chlorides of the metals, exactly instances in which combinations are formed by them with chloride acid. Such compounds, however, are rare, and, when they do occur.

pear not to possess stability.

by Berzehus as fluchydroboric acid and fluchydrosiheic acid, should be considered as tertium quids, in which the fluoride of hydrogen per at the part of a base, while the fluorides of boron and silicon act as and Hence fluohydroboric acid is a fluoborate of hydrogen, and fluohydrobacid, a fluosilicate of hydrogen.

hydrogen forms compounds analogous to these abovementioned, and visit would designate as fluocolumbate, and fluotitanists of hydrogen.

2083. When any fluosalt like those abovementioned, in which hydrogen exists as a radical, is brought into contact with an oxybase, of which the radical is capable of forming a fluobase, the hydrogen unites with the stygen of the oxybase, while the radical of this base unites with the fluores. The fluocid of the fluosalt, consequently liberated, combines with the resulting fluobase.

20%4. The other fluoralts are formed by the direct reaction of the fluorals and fluobases which compose them, either in the wet or dry way. By adding fluohydric acid to the fluorides of potassium and sodium, fluohydrass

of those thobases may be formed. (1398.)

SECTION V.

OF CYANOSALTS.

2085. The cyanosalts are in general obtained either by the direct are of a cyanocid with a cyanobase, or by decomposition. It is by the late method that the cyanoferrite of potassium is formed, the sulphase of the protoxide of iron being presented to the cyanide of potassium. In this case the sulphane acid, and the oxygen of the protoxide of iron, are transferd to one portion of the potassium. The cyanogen, consequently therest forms with the iron, cyanoferrous acid, which unites with the underposed portion of the cyanide of potassium. (1299, &c.)





COMPENDIUM

07

THE COURSE OF CHEMICAL INSTRUCTION

IN

THE MEDICAL DEPARTMENT

OF

THE UNIVERSITY OF PENNSYLVANIA.

BY

ROBERT HARE, M.D.

PROFESSOR OF CHEMISTRY.

PART II.

COMPRISING THE

CHEMISTRY OF ORGANIC SUBSTANCES; BEING A COMPENDIOUS SELECTION FROM THE PREVIOUS EDITION: THE "TREATISE OF ORGANIC CHEMISTRY," BY LIEBIG: GREGORY'S TURNER: KANE'S "ELEMENTS," AND THOSE OF GRAHAM: INTERSPERSED WITH SOME ORIGINAL MATTER.

Also, a Letter on the Berzelian Nomenclature, with the Reply of Berzelius; with some Subsequent Remarks and Suggestions by the Author.

And an Effort to Refute the Arguments advanced in favour of the Existence of Compound Radicals, like Cyanogen, in the Amphide Salts; with a subjoined Statement of those Arguments as advanced in Kane's Elements.

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### OF ORGANIC CHEMISTRY,

OR

### THE CHEMISTRY OF ORGANIC SUBSTANCES.

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# OF ORGANIC CHEMISTRY,

OR THE

### HEMISTRY OF ORGANIC SUBSTANCES.

5. Under the appellation of organic substances are ised—

7. 1st. All those which are created in vegetables limals.

3. 2dly. Such as are generated from those above ned, either by spontaneous changes, aided by temre or catalysis, or by reciprocal reaction.

). 3dly. Such as arise from the substances created erated as above described, in consequence of their

on with inorganic bodies.

- D. In this department of the science it is, perhaps, less t to acquire some general ideas, than to make an progress in the chemistry of inorganic substances, timate elements of vegetable and animal matter are and are peculiarly well known. But the light which we upon inorganic compounds, by resolving them seir ultimate elements, is much more satisfactory ny which we can, by the same means, extend to orproducts. Between these, ultimate analysis can deate little more than a difference in the proportions hydrogen, oxygen, carbon, and nitrogen, of which re constituted; although in their influence on vitality may display the opposite properties of the most defood, or the most deleterious poison; of delighting nding our senses in the extreme.
- 1. Hydrogen, oxygen, and carbon, are the principal te elements of vegetable substances; especially carhich is pre-eminently essential to their constitution, as been alleged to perform, in vegetables and aniapart analogous to that which silicon performs in ils.
- 2. In some essential oils, in caoutchouc, in ammonia

in cyanogen, and in some compounds formed or derived from these substances, there is no oxygen, while in oxaic acid, and some other oxides of carbon, no hydrogen exists. But in no instance, excepting that of ammonia, and its hypothetical associates amide and ammonium, is carbon deficient; and in the great majority of instances, the three elements above named are indispensable ingredients. Although, comparatively, nitrogen be found only in a few substances, those into which it does enter are generally pre-eminently active in their properties; and, agreeably to Liebig, without its assistance, vegetation cannot three Hence, as he alleges, it is always to be found in vegetable organs, although not a constituent of many substances which they secrete or excrete.

2093. Magnesium, calcium, sulphur, phosphorus, incessilicon, bromine, iodine, fluorine, are also found in minute proportions in certain parts of certain vegetable or animal products; and it may be inferred that they perform some useful office; but although subservient, in an important degree, to the functions of animals and plants, they are constituents neither of their organic tissues, nor secretory products.

2094. It is generally a marked distinction, between organic and inorganic products, that the latter can, in a much greater number of instances, be imitated by art.

2095. The incompetency of chemists to regenerate the substances analyzed by them, has caused the accuracy of their deductions to be questioned. Rousseau, having heard Rouelle lecture on farinaceous matter, said he would not confide in any analysis of it, till corroborated by its reproduction from the elements into which it was alleged to have been resolved. I conceive that an acquaintage with facts, thoroughly demonstrable by modern chemistry, would have rendered that ingenious philosopher less seeptical. At first view it may seem reasonable to consider synthesis as the only satisfactory test of the truth of andlysis. But if when diamond is burned in one bell glass, and charcoal in another, in different portions of the same oxygen gas, and subsequently, in each vessel, in lieu of the diamond and charcoal, carbonic acid is found, from which by potassium, carbon may be liberated, who would hertate to admit both substances to consist of carbon, because this element cannot be recovered in its crystalline form from the gaseous state?

# Of Organic Hydrates.

2096. It was suggested by Prout, that as, in many vegele substances, consisting only of carbon, hydrogen and gen, the two last mentioned elements existed exactly in proportion for forming water, they might be consied as constituted of water and carbon, and, consently, as hydrates of carbon.

2097. But it has since been shown, that either the hygen of these supposed hydrates, may, in various inces, be supplanted by other elements without separathe oxygen; or that the oxygen may be supplanted

hout separating the hydrogen.

1093. It is, however, an important and interesting fact, t almost all vegetable substances which are neither 1, oily, nor resinous; such, for instance, as gum, sugar, ch, lignin, hold the elements of water in the ratio resite to form this liquid, however these elements may be sciated.

the Influence of Heat upon Vegetables, with and without Access of Air.

1099. When subjected to distillation, vegetable subices devoid of nitrogen, in the first place, yield the water essential oils previously existing in them. At a higher perature, certain essential oils or spirits, analogous to shol, resulting from a new arrangement of the ultimate nents, are in some instances evolved; and either at the ne time, or subsequently, at a higher temperature, acetic 1, associated with bituminous or empyreumatic matter, h carbonic oxide or carbonic acid, and carburetted hygen are generated. By further ignition, the volatile ducts thus obtained, may be resolved into carbonic oxand carburetted hydrogen; a deposition of carbon he solid or pulverulent state, being always a concomit of the change. In proportion as the hydrogen is raed by heat, its capacity to suspend the carbon appears se diminished (1259). So far as nitrogen is present, the union of an atom of this element with carbon or lrogen, ammonia or cyanogen, or some of their comnds, may be generated.

2000. The results above mentioned evidently proceed, a great measure, from the superior volatility of the

hydrogen and oxygen, which causes them to pass off into the aëriform state, with such portions of the carbon as they may, under these circumstances, be capable of retaining.

3001. The experiments of Sir James Hall show, that we getable matter, wood for instance, when subjected to have and pressure, is converted into a bitumen analogous to that of mineral coal. Under these circumstances, caloric destroys the organic structure, but does not sever the constituents of many bodies, which would be otherwise partially dissipated. When ignited in the air, it were almost unnecessary to say that hydrogen, oxygen, and carbon must yield water and carbonic acid only. There are the only products of hydrogen and carbon, when burned where there is an ample supply of oxygen.

3002. By a carefully managed heat several vegetable acids may be converted into acids of a different kind. In some instances, difference of temperature is sufficient to

vary the character of the resulting acid.

# Of the Ultimate Analysis of Organic Substances.

been latterly accomplished by heating the substance with oxide of copper, so as to oxidize all the carbon and hydrogen, and liberate, in the gaseous state, any nitrogen which may be present. The hydrogen has been in general estimated from the water produced; the carbon, from the quantity of carbonic acid. Hence the products of the operation have been first passed over chloride of calcium, and afterwards subjected to hydrate of potash, lime-water, or alkaline solutions. The water is estimated from the increased weight of the chloride, and the carbonic acid by the volume absorbed, or the increased weight of the alkaline solution employed for its detention.

3004. By Messrs. Will and Varrentrapp, the proportion of nitrogen in a compound has lately been ascertained by heating it with a mixture of quicklime and hydrate of soda, in a tube of refractory glass. Under these circumstances, the element in question, uniting with hydrogen to form ammonia, is easily secured by means of a dilute solution of chlorohydric acid. The resulting chloride of ammonium is precipitated by chloroplatinic acid, and the resulting salt is washed in a mixture of ether and alcohol. The quantity of nitrogen is estimated by the table of equi-

and by ascertaining the loss of weight consequent sure to a red heat. Agreeably to the table of

ents, 16 of the loss thus sustained is nitrogen.

. When chlorine is present, chromate of lead is lieu of the oxide of copper, because the chloride of being volatile, would be carried into the cavities ed for the absorption of water and carbonic acid.

. When liquids are to be analysed, small portions oduced into glass bulbs so as to alternate in a tube ide of copper, or some other oxidizing agent.

### Mode in which the Ultimate Ponderable Elements of Organic Bodies are associated.

- . As in the analysis of the mineral kingdom, we deas elementary, those substances which we cannot further, so, in examining organic products, those ces of which the grouping cannot be altered without ing their most important characteristics, are to be as the elementary principles, by which the nature pounds is to be understood and described.
- . Liebig alleges, that the principal object of organic try, is the investigation of the properties and comn of organic combinations, and the mode in which ements are grouped. The idea attached to the word l in this instance, may be illustrated by contempla-: formula of a compound in one way, so as to exhithe proportions in which each ultimate element n it; in another way, so as to make evident not eir proportions, but their grouping likewise. mula C<sup>2</sup> O<sup>3</sup> shows, that two atoms of carbon and f oxygen enter into the composition of oxalic acid;

× CO<sup>2</sup> shows, that this acid is composed of car-

xide CO, and carbonic acid CO<sup>2</sup> (556, &c.).

. In like manner, cyanhydric acid may be repreas a compound of two atoms of carbon, one of niand one of hydrogen, C<sup>2</sup> N H, or as a compound ogen, C<sup>2</sup>N and hydrogen, H, formula, C<sup>2</sup>N+H.

The compounds thus cited, CO carbonic oxide, and

anogen, are considered as acting as compound ra-

This appellation is employed to designate in these es and in others, certain groups of ultimate elewhich appeared to be endowed with the power, like simple ultimate elementary atoms, of entering in combination with one or more of their composing atoms

or of other simple elementary or compound atoms.\*
3011. From a deficiency of better words I shall considerate. a "compound radical," so called by Liebig, as a compound element, when, like cyanogen or ethyl, it acts as a simple element. I shall restrict the use of the name ration agreeably to the definition in my Inorganic Chemistry, such bodies as do not form the common ingredient of acid and a base.

3012. Compound elements, like cyanogen, which, they unite with an anion and a cathion, form with former an acid, with the latter a base, I consider as be-

longing to the basacigen class (627).

3013. As on the one hand, it is seen that cyanogen forms the part of a basacigen body, or one capable of ducing acids and bases, by combining with radicals; so, the other, we may perceive ammonium, consisting of the drogen and nitrogen, N × H<sup>4</sup>, capable like a metallic radical of forming compounds with the basacigen class, which have basic properties in some instances of great energy But latterly, pursuant to the suggestion of Kane, ammon is conceived to contain a compound element analogous cyanogen, consisting of NH2 which is called amide, and which combines with hydrogen and other radicals, forming compounds called amidurets, capable of union with other definite compounds. Thus it is inferred, that white precipitate consists of amide, mercury and chlorine, NH' Hg + Cl Hg, the symbol of amide is Ad, which being substituted in the above, we have Ad Hg + Cl Hg for the formula of white precipitate.†

3014. This view of the subject is now generally sanctioned, although neither amide nor ammonium have been

isolated.

3015. In fact, it has been shown of late, that there are a

t N is the symbol of nitrogen, H of hydrogen, Cl of chlorine, Ad of amide, Hg of

hydrargyrum or mercury (556, &c.).

<sup>\*</sup> Strictly, an element cannot be compound; but chemists, before the idea of compound radicals originated, distinguished compounds capable of entering into bination and of being separated again, and transferred to other compounds, as premate elements, in contradiction to simple elements also called ultimate elements Upon this view of the subject, the ultimate analysis has been understood to comp the idea of the resolution of a substance into its simple elements, in contradisting tion to an analysis by which its proximate elements are separated. Alcohol 🕪 jected to ultimate analysis would be converted into hydrogen, oxygen and carbon while by another procedure, it may be resolved into its proximate elements with and ether. I feel myself authorized, under this view, to call those bodies composed elements, which, consisting of more than one element, act like simple elements.

table or animal matter, as capable of uniting with bodies as do elementary radicals, forming like des, chlorides, bromides, iodides, fluorides, cyanhides, &c. Of the compounds thus produced, the part of a radical in an acid, some an analoe in a base or even of an alkaline base. Morecids and bases thus produced, unite similarly to erated by a union of ultimate elements, which n many cases competent to displace from com-

compound organic radicals may be divided into ses accordingly, as capable of forming acids, or neither. Hence, they may be distinguished as, as basifiable, or as indifferent.

The acidifiable compound radicals are as follows:

| oxide or pro<br>or bicarbur<br>sesquicarbu | -<br>- | Formula. CO C3 N C6 N4 |   |   |                                               |
|--------------------------------------------|--------|------------------------|---|---|-----------------------------------------------|
| benzule or b                               |        | -                      | - | - | C14 H5 O2                                     |
| or cinnamul                                |        | -                      | - | - | C <sub>18</sub> H <sub>8</sub> O <sub>2</sub> |
| · salicule,                                | •      | •                      | - | • | C14 H5 O4                                     |
| acetule,                                   | -      | -                      | - | - | $C_4$ $H_3$                                   |
| · formule,                                 | -      | -                      | - | - | $C_3$ H                                       |

The basifiable compound radicals are

|            | -   | - | - | - | - | $N H_3$                       |
|------------|-----|---|---|---|---|-------------------------------|
| thule,     | •   | - | - | - | - | C4 H5                         |
| methule    | , - | • | - | - | • | $C_3$ $H_3$                   |
| etule,     | -   | • | - | • | - | C35 H33                       |
| or glyceri | ule | • | • | - | - | C <sup>6</sup> H <sup>7</sup> |
| .mule      | -   | - | - | • | - | $C^{10} H^{11}$               |
| ·misityle  | ne  | - | - | - | - | C <sup>6</sup> H <sup>4</sup> |
| r kacodu   | ıle | • | • | • | - | C4 H6                         |

There are likewise some subordinate compound

swith very few exceptions in formulæ expressing sition of organic substances, only four different requisite, with the figures showing the relative s, the employment of symbols for that purpose y highly advantageous. The student, therefore,

resolution, any repugnance to the study of the formula above given, or others which may be resorted to in the or in other modern treatises of chemistry. A compansat of their formulæ, respectively, will convey an idea of the difference in composition existing between the radicals in the preceding list.

3021. Agreeably to Liebig, the term "compound radical denotes a class of compound bodies possessing the capacity of uniting with the simple elements, and forming, with them, combinations analogous in their properties to con-

binations of two simple elementary bodies.

3022. From combinations formed as above mentional the simple element may be removed and replaced by another element, simple or compound.

3023. According to the same authority, compound radicals are capable of combining with each other, and

forming acids with oxygen, sulphur, or hydrogen.

arranged in groups, each derived from their appropriate compound radical by the combination of this radical with elementary atoms, and the union of the resulting com-

pounds with other compound bodies.

analogy in composition. It depends exclusively on the composition. It depends exclusively on the composition. It depends exclusively on the composition.

3026. In consonance with the law in question, Dunes has found, that in acetic acid chlorine may be substituted for hydrogen, and that in this way a new acid, designated

as chloroacetic, may be produced.

3027. This chloroacetic acid is by him alleged to be, in

perties, so analogous to acetic acid, that to know abitudes of the one, conveys an idea of those of the

This analogy he conceives to arise from a chemiw, agreeably to which the properties of a compound I rather on the type of the composition, than on the ular character of the elements which may have been nged.

- 8. Berzelius asserts that chloroacetic acid differs from acetic acid in properties, and that the facts adjustify nothing beyond an opinion, originally exdupon the subject by Dumas himself, who, speaking law of substitution, admitted it to be an "empirical eserving our attention only so long as it might hold
- 9. It appears to me, that the facts of the cases adto in the support of the doctrine of substitution, strate them to come under the fourth case of affi-523), in which two bodies, simple or compound, in union, another body, added in excess, unites with

O. In the case of acetic acid exposed to an excess of ne, there is the affinity between hydrogen and chlound that between chlorine and the elements, with

hydrogen is previously combined.

1. Hence results chlorohydric and a new acid, called acetic, in which chlorine may act as a radical, as it is to do in its combinations with oxygen. The exist-f chlorocarbonic acid demonstrates that the display nity between chlorine and oxides of carbon, is not an ily.

2. Either of the classes of radicals abovementioned, e distinguished into primitive and derived radicals.

1, a sesquicarburet of nitrogen, is derived from cyn; and acetyl and formyl, from ethyl and methyl.

3. Being convinced that in the present state of che, more even than heretofore, it is best to aim at
il knowledge first, and afterwards to proceed to
ulars, I shall not treat of the compounds formed
adicals or products obtained from them, under their
respectively, unless where the substances alluded to
practical importance.

19

### Of Amide, NH2.

3034. Ammonia, it will be remembered, consists of atom of nitrogen and three of hydrogen, NH. assumed to consist of one atom of nitrogen and two

hydrogen, as the formula above given indicates.

3035. The phenomena which ensue when potassium i heated in ammonia, had long been an object of unsucce ful speculation. The metal, when so exposed, become converted into an olive-coloured mass, which, by coak with water, is converted into potash and ammonia.

3036. I believe that Dr. Kane was the first to suggest that in this case the alkalifiable metal takes the compound radical, amide, from the ammonia. Thus a compound generated of amide and potassium. When the amider of potassium, produced as described, is presented to water, this liquid regenerates ammonia, by supplying 🗮 additional atom of hydrogen to the amide, while the potassium is, by simultaneous oxidizement, converted in potash. It follows that ammonia is an amiduret of by drogen.

3037. Compounds of amide are called amides by Liebing amidides by Kane; though it will be seen, that when combined with hydrogen, Liebig designates the resulting com pound as a hydruret; in French, hydrure. Consistently with the nomenclature which I have employed, the termination in ide is restricted to the basacigen class; I shall therefore use the termination in urci for the compounds of amide. It is singular that Liebig should use the same

word, amide, for the radical and for its compounds.

3038. As by the subtraction of an atom of hydrogen from ammonia, amide is generated, so, by the addition of a like atom, we generate ammonium, of which I have al-

ready treated (1106).

3039. Liebig represents amide as acting with hydrogen in the place of an electro-positive radical. Hence, agree ably to his language, ammonia is a "hydrure d'amide," is English, a hydruret of amide; or, more briefly, he calls it hydramide. Of course, consistently, ammonium is a bibydruret, or bihydramide.

3040. The following formula will serve to explain the composition of some of the compounds of amide. It should be kept in mind that Ad is the symbol of amide.

recipitate is a compound of amiduret of mercury and bihloride, Ad, Hg + Hg Cl<sup>2</sup>. Another amido-chloride is rmed by the reaction of white precipitate with alkalies, hen results a compound of an amiduret with the bioxide and bichloride of mercury, Ad Hg + Hg O<sup>2</sup> + Hg Cl<sup>2</sup>. wo atoms of amiduret of mercury unite with a subsulhate, whence we have a biamido-subsulphate, Ad Hg<sup>2</sup> + o<sup>2</sup> 2HgO. Biamido-sesquinitrate of mercury consists of vo atoms of amiduret of mercury, two of acid, with three f bioxide of the same metal, 2Ad Hg + 2NO<sup>5</sup> 3Hg O<sup>2</sup>. mido-subnitrate, consisting of an atom of amiduret and vo of subnitrate, Ad Hg NO<sup>5</sup> 2HgO.

3041. White precipitate has been designated as chlormide of mercury: I prefer the name, above employed,

f amido-chloride.

# Of Carbonic Oxide as a Compound Radical.

3042. Of this radical I have already treated as an oxide f carbon. By combining with carbonic acid, CO<sup>2</sup>, it contitutes oxalic acid, C<sup>2</sup> O<sup>3</sup>. This acid, by combining with ydrated ammonia, NH<sup>3</sup> + HO, or more properly with oxide of ammonium, consisting of the same ultimate elements ifferently grouped (1116), forms neutral oxalate of ammonia, so called. This oxalate is principally used as a test or lime.

3043. Chloroxycarbonic acid is a product of the union of arbonic oxide with chlorine, of which some mention has

een made (1240).

3044. Carbamide is the name given to a compound ormed by the union of carbonic acid with amide. It is btained by mingling chloroxycarbonic acid with ammonia. In this way solid white crystals are produced, conisting of carbamide and chloride of ammonium.

3045. Oxamide, C<sup>2</sup> O<sup>2</sup> + Ad, consists, as may be seen rom the preceding formula, of two atoms of carbonic oxde and one of amide. It may be designated as an amido-

oxide of carbon.

3046. This compound is obtained in great purity, by lecomposing oxalic ether by liquid ammonia, or by heating an oxalate of ammonia in a retort, with a receiver annexed. The oxamide passes into the receiver, and conlenses in white flocks. These, being insoluble in water, are depurated by washing with this liquid upon a filter.

3047. Oxamide is described as a brilliant white powder, insoluble in alcohol, ether, and in cold water, but soluble, in a small proportion, in hot water.

3048. Subjected to dry distillation, it is resolved into water, carbonic acid, cyanhydric acid, cyanic acid, and

ammonia.

3049. Oxamide differs from the common oxalate of ammonia, consisting of oxalic acid and oxide of ammonium,

in having two atoms less water.

3050. Oxalic ether, which may be decomposed instantly by an aqueous solution of ammonia, consists of anhydrous oxalic acid and ether, or oxide of ethyl. The acid yields an atom of oxygen to one of hydrogen to form water, by which the ether is converted into alcohol, while on the one side there remains carbonic oxide, CO, on the other, amide, NH<sup>2</sup>, which by reciprocal union constitute oxamide.

3051. When examide is heated with alkalies or acids, by the accession of an atom of water, exalic acid and ammonia are generated. The same result ensues from the exposure of a mixture of examide and water, to a tempera-

ture above the boiling point.

### Of Benzule, & Ol zoile or Benzyle, CH, Hs, O2.

3052. The preceding name is given to a compound radical inferred to exist in benzoic acid, and in the essential oil of bitter almonds, giving rise to several interesting compounds. By the addition of an atom of oxygen and an atom of water, it forms crystallizable benzoic acid, which like many other acids, cannot exist without an atom of water, or some other base.

3053. By the substitution of an atom of hydrogen for an atom of oxygen, benzoic acid is converted into the pure essential oil of bitter almonds, C<sup>14</sup>, H<sup>3</sup>, O<sup>2</sup>, which Liebig

designates as a hydruret of benzule.

3054. By bringing either of the halogen bodies (627), or various acids in contact with this hydruret, with or without exposure to the distillatory process, a variety of compounds may be produced. These compounds, in composition and properties, are somewhat analogous to ethers; inasmuch, as they mix either with ether or alcohol, and retain their radical with an energetic affinity.

3055. The hydruret of benzule does not pre-exist in bitter almonds, but is the product of the mysterious catalyzing influence of two substances which they contain, amigdalin and emulsin, or synaptase, in an aqueous mixture when subjected to distillation. During the reaction thus induced, cyanhydric acid being generated, endows the resulting oil or hydruret, with a well known poisonous property, which, in the absence of that acid, has been ascertained not to exist.

3036. Benzule forms a compound with amide, called benzamide, by the reaction of chloride of benzule with dry ammoniacal gas; and likewise an acid, by uniting with formic acid, called formobenzulic acid. Hippuric acid, which is the uric acid of the horse, consists probably of benzamide and another peculiar acid; or of hydruret of benzule, with cyanhydric and formic acids.

3057. According to Mr. Alexander Ure, benzoic acid taken internally by man, is discharged in the urine as hippuric acid, the proportion of uric acid undergoing a cor-

responding diminution.

# Cinnamyl, C18, H8, O2.

3058. Between this radical and benzule, there is much analogy; since cinnamyl plays a part in pure oil of cinnamon, or hydruret of cinnamyl and connamic acid, analogous to that which benzule plays in its hydruret and in benzule acid. In either case, the substitution of oxygen for hydrogen, converts the hydruret into an acid having the same radical.

3059. Cinnamyl exists in oil of cinnamon, which, when pure, constitutes a hydruret, and in an acid called cynnamic, playing a part similar to that which benzule has been represented as performing in two analogous compounds. In either case, the oily hydruret may be converted into an acid having the same radical, by the substitution of an atom of oxygen for one of hydrogen.

3060. This radical is said to exist in an oil, separable

from the balsams of Peru and Tolu.

3061. It does not appear that the compounds formed

with this radical are numerous or important.

3062. By the reaction of pure colourless nitric acid with Chinese oil of cinnamon, a crystallized nitrated hydruret of cinnamyl may be obtained, C,  $H^8$ ,  $O^2 + HO + NO^5$ , which by addition of water liberates the pure hydruret of cinnamyle,  $C^{18}$ ,  $H^8$ ,  $O^2$ .

### Of Salicyl, C14, H5, O4.

3063. This hypothetical radical is inferred to exist in the oil of the spirea ulmaria or queen of the meadow, and in that evolved from a neutral crystallizable substance, called salicin, which may be extricated from the leaves and bark of any species of willow, of which those parts have a bitter taste; and also from some species of poplar. It was originally discovered by Buchner and Lerouz, in the bark of the salix helix.

3064. This radical has a great analogy to benzule in properties, as well as proximity in composition, as must be evident from a comparison of their respective formula.

It is in fact benzule plus two atoms of oxygen-

3065. The oil above mentioned has the same relation to salicyl, that the oil of bitter almonds has to benzula both being hydrurets. The oil of spirea is treated as a hydracid, which it will be well to keep in mind when sourness is insisted upon, as a property peculiar to what are improperly called hydracids.

3066. This hydruret may be obtained in a state which is isomeric, if not identical with that in which it is extricated from the spirea, by distilling one part of salicin with three parts of bichromate of potash, four and a half of sulphure

acid, and thirty parts of water.

3067. It is a colourless, oily, inflammable liquid, with a burning taste; density 1.731, freezing at 4°, boiling at 335.7 when obtained from spirea, but 359.6 as obtained from salicin. It dissolves easily in water, alcohol and ether.

3068. Salicyl, like benzule, forms compounds with the halogen bodies, or with acids, by their reaction with its hydruret. The reaction with ammonia differs from that of benzule, as it unites with the ultimate element, nitrogen, instead of amide.

### Of Ethyl, C4 H5.

3069. If this be really, as is generally now believed by chemists, the radical in the well known liquid, alcohol certainly, for good or evil, it is one of the most important and interesting compounds in nature.

3070. In the year 1836, in the last edition of my Organic Chemistry, agreeably to the doctrine prevailing at the time, I treated alcohol as a compound of two atoms of

r and one of etherine, C<sup>4</sup> H<sup>4</sup> + 2HO. Common ether ring from alcohol only in having an atom less of r essential to its constitution, was represented to be prohydrate of etherine, C<sup>4</sup> H<sup>4</sup> + HO. No change has a place as to the ultimate analysis of these liquids.

only as to the grouping of the ultimate elemenconstituents, by which we have in ether, C<sup>4</sup> H<sup>5</sup>O, in alcohol the formula of ether, with an additional of water, C<sup>4</sup> H<sup>5</sup>O + HO. Thus, instead of a monoate of etherine, ether becomes an oxide of ethyl; and nol, from a bihydrate of etherine, is transferred into a ated oxide of ethyl.

71. Agreeably to either view, the transformation of nol into ether requires only the removal of an atom of r.

- 772. It is well known that common ether may be obd by the distillation of alcohol with sulphuric acid, that when, to the materials employed for this purpose, cid, or a salt containing an acid, is added, an etherial bound of ether with the acid, having more or less anawith common ether in properties, may be obtained.
- r bearing its name has not been formed, such as nitric r, acetic ether, tartaric ether, oxalic ether, muriatic r, &c.
- 74. The rationale is evident: as to convert alcohol ether, the removal of an atom of water is all that is isite; to generate any other ether, it is only necessary this oxide, in its nascent state, shall be in contact with icid, or be presented to any basacigen body in union hydrogen; so that the ethyl may be deoxidized by ormation of water, and presented naked to the basacielement.
- 75. Under this view of the composition of ether, it naccountable, that this oxide will not combine with , excepting when it is in a nascent state; but this obn may apply also to the existence of etherine as the of the ethers.
- 76. It does not appear that ethyl has ever been iso-L. I have not only distilled pure ether from potas-, without decomposing it, but have likewise cohobated ith potassium in a glass tube, hermetically sealed. lower end, to which the contents naturally subsided,

was kept boiling by a water bath for several days, without being decomposed more than partially. The potassium became coated with a white crust, which being removed, the metal appeared in its metallic state.

3077. The etherial compounds of cthyl may be class-

fied as forming one order of ethyl ethers.

3078. We have then, in this order, the following class-es:—Class 1st. Simple ethers, formed by the union of chyl with any basacigen element which are named after such element.

3079. In this class we have the

Oxide Chloride Bromide Iodide Sulphide Selenide Telluride

of ethyl.

3080. Complex ethers are formed by the union of as acid with any one of these. Excepting those formed with the oxacids and sulphydric acid, there are no ethers in this class. The oxacid ethers may be considered as forming a genus comprising an etherial compound for almost every acid of importance.

3081. There is only one sulphacid ether, mercaptan, or

the sulphydrate of the sulphide of ethyl-

3082. In consequence of its being obtained by the distillation of sulphuric acid with alcohol, the oxide of ethyl was formerly called sulphuric ether, and is still mentioned under that name in commerce, agreeably to the opinion that it consisted of water and etherine, as other ethers consisted of etherine and an appropriate acid. In the list edition of my Organic Chemistry, I designated this oxide as hydric ether. It is a curious consequence of the change which has taken place, as above described, in the prevailing opinion on this subject, that the name above mentioned is now due to alcohol, which, as respects composition, is, in fact, hydric ether. Yet it differs from ethers in general in having a strong affinity for water in all proportions.

3083. It may be well to premise, that I shall adopt for the oxide of ethyl, when not particularly desirous to recall its chemical composition, the usual name of other, which

ly claim by prescription, however temporarily it may

been otherwise designated.

84. As alcohol differs from ether only in the presence atom of water, it follows that any chemical reaction, he should effect the removal of that atom, ought to ert it into ether. Yet, excepting the reaction, during lation, with one or two chlorides, a resort to which d not be found economical, the conversion of one liinto the other is accomplished by a most complicated intricate play of affinities, which has been a most prosource of discussion among the most eminent ches. Nevertheless, this subject is still debateable, notstanding that much light has been thrown upon the acpanying phenomena.

185. It may be well for the student to recollect the ive composition of these important liquids, and that conversion of one into the other arises from the subon of alcohol, mingled with certain acids, to the dis-

ory process.

86. When sulphuric acid is employed as is usual, the result is a combination between two atoms of this one of oxide of ethyl, and one of water, forming what been called, heretofore, sulphovinic acid, or what Lielesignates as the acid sulphate of the oxide of ethyl.

87. Evidently it would be more properly defined as a le sulphate of ether and water;\* for as, what is called entrated sulphuric acid, when deprived of water as far is effect can be produced by ebullition, is a sulphate ater; sulphovinic acid, consisting of one atom of this late, and one atom of sulphate of the oxide of ethyl, be a double sulphate of the oxide of ethyl, and wa-

# 88. So long as the proportion of water present in

so that when a metal, by contact with the acid, displaces hydrogen, it is merese in which one radical supplants another. Agreeably to a new doctrine, all phur and oxygen present, acts as a compound radical, and, as such, is transfrom one radical to another; but this I think I have shown to be untenable. Fort to refute the Arguments in favour of the existence of Compound Radicals phide Salts, 6, 92.

order to understand the above given explanation, it should be recollected, so beiling point of diluted sulphuric acid rises, as the proportion of water in with it lessens, till it attains the point at which the sulphate of water itself ses, which is about 600°: also, that the affinity of concentrated sulphuric acid ser is so great, as to enable it to abstract the elements of this liquid from orsubstances; in which case they are blackened, and said to be carbonized, in

nence of the evolution of carbon.

the mixture of sulphuric acid and alcohol, is adequate to keep the temperature sufficiently low, the ether, in the double sulphate, being more volatile than the water, existing in excess in the solution, yields the acid to this equid, and comes over, accompanied by a proportional quantity of steam, and at the outset, of alcoholic vapour. Thus ether, alcohol, and water, being partially removed, the proportion of acid relatively to the residual materials, is uncreased: but as this takes place, its avidity for water augments, and the boiling point of the mixture rises. In consequence of the increased avidity for water, the acid takes from a portion of the ether, C' H'O, an atom of each of the elements of this liquid, HO. Thus etherine s

evolved, C' H'.

3089. Meanwhile the increased heat causes a portion of the etherine to give up the whole of its hydrogen to 1 part of the oxygen, of a portion of the acid. Hence suphurous acid and carbon are evolved; the one being indicated by the carbonaceous colour, the other by its well known suffocating fumes. Under these circumstances, a triple compound, consisting of sulphuric acid and oxide of ethyl, and a portion of undecomposed etherine, being formed, comes over with sulphurous acid and ether, forming a yellow liquid. When this liquid is deprived of its sulphurous acid by ammonia, or any other alkaline base, and the ether is removed by distillation, the residue is the liquid long known as oil of wine, being the efficient and characteristic ingredient of Hoffman's anodyne, erroneously represented in several European works as a more mixture of alcohol and ether. The preferable mode to isolate the of of wine, is to expose the yellow liquid, in vacuo, over subphuric acid in one capsule and slaked lime in another. The sulphurous acid is absorbed by the lime, the ether by the sulphuric acid. The quantity of acid in the oil varies with the mode of isolation; being greatest when the last mentioned mode is resorted to.

3090. The word ether was originally employed to designate a supposed elastic aëriform matter, vastly more ran and subtile than air. It is still used in that sense as an appellation for the matter, which is, according to the undulatory theory, the medium by which luminous bodies radiate light. By analogy, the word other was employed to designate a liquid which bore the same relation to other

iquids, as ether proper to air. This appellation has naturally been extended to other liquids analogous in properties and composition. Of ethers in general, common ether nay be considered as the best exemplification. What nainly distinguishes the liquids thus called, from alcohol, is their very inferior miscibility with water. Many of them are, however, heavier than water, so that, upon the score of density, they do not merit to be distinguished as etherial.

3091. It will be seen that there are several hydrates, ormed with other compound radicals, which are congeners of alcohol in composition, and, to a limited extent, resem-

ole it in properties.

3092. Generally, substances considered as etherial are susceptible of distillation, are inflammable, little soluble in water, but highly susceptible of union with alcohol, essential oils, and resins. They are, for the most part, fragrant and stimulating to the taste, affecting the animal nerves cowerfully when inhaled, or swallowed, even in a minute quantity.

# Of Acetyl, C<sup>4</sup> H<sup>3</sup>.

3093. The preceding name has been given to a hypochetical sub radical containing the same number of atoms of carbon as ethyl, with three atoms of hydrogen instead of five. This radical is inferred to play the same part, in liquid lately discovered and called aldehyde, that ethyl loes in alcohol. In fact, the only difference in composition existing between these liquids, is that between their radicals; the former being produced from the latter by the emoval of two atoms of hydrogen.

3094. Acetyl is chiefly interesting as the radical of the mportant acid of vinegar, designated by modern chemists acetic acid. This acid, in the hydrated state, in which done it is capable of isolation, results from the addition of wo atoms of oxygen to aldehyde. By the lesser addition of one atom of the same element, another acid has been

made, called acetous acid, or aldehydic acid.\*

As both this acid and acetic acid have the same radical, the compound, having he lesser proportion of oxygen, should terminate in ous (1052, &c.). Hence the cid in question should be called aldehydous acid, if named, so as to show its derivation from aldehyde, and acetic acid should be called aldehydic acid; but aldehyde teelf enters, as an acid, into an ammoniacal compound, the formation of which is a recursory step in obtaining it in a state of purity. Of course, if these compounds

3095. By Liebig, olefiant gas is treated as a hydrust of acetyl, C' H3 + H = C' H4, which is just double the quantity of carbon and hydrogen contained in a volume of olefiant gas. But, according to Berzelius, the two atoms of carbon, and two atoms of hydrogen, in a volume of this gas, constitute an independent radical, which he calls clayl. Agreeably to Liebig's view, olefant gas is isomeric with etherine, or etherole, the name given to etherine by him.

3096. Agreeably to the view of the former, the oil resulting from the reaction of olefiant gas with chlorine, is a chlorohydrate of chloride of acetyl, C' H' Cl + HCl, while, if the Berzelian idea be adopted, it consists of two atoms

of elayl and two of chlorine, C' H' Cl2.

3097. When this compound is dissolved in a solution of potash and alcohol, it is decomposed into chlorohydric acid, which forms water and chloride of potassium with the potash, and a compound, which escapes in the gaseous form, consisting of C' H3 Cl. The composition of thus gas is evidently such, that it may be considered as a chloride of acetyl; and its formation must be regarded as confirming the view taken by Liebig of the composition of the oil of the Dutch chemists. Bromine, like chlorine, on being presented to olefiant gas, produces a compound, which may either be considered as a bromohydrate of the bromide of acetyl, or simply as a bromide of clayl, in other words, of olefiant gas; but which, by reaction with the alkalies, evolves a gas, the composition of which, it would seem, can only be reconciled with the idea of a bromide of acetyl. The action of iodine is analogous, but not so well ascertained. The product is pulverulent in its consistency, but in other respects resembles that which results from the reaction of chlorine with olefiant gas.

### Of Mesityl or Misitylene,\* C6 H4.

3098. The vapour of pure acetic acid, in passing through a red-hot porcelain tube, is decomposed, yielding a colour-

be all considered as exacids of acetyl, as I think would be more proper, agreeably

the nomenclature adopted in analogous instances, their names would be accept and accepts acid. But aldehyde, as a congener of alcohol, is, purhaps, preferably designated as a hydrated exide of acetyl.

Liebig does not introduce this radical into his general list of radicals, but trees of it as a product of the decomposition of acetyl. The same course is pursued a respect to hacodyl, although this contains, as will soon appear, arresis, an element which does not axist in acetyl. It will also be found that he places august under ethyl, as yielding ethyl by their decomposition. This does not appear to me judicious, because, by the same rule that mosityl is placed under acetyl, ethyl abould seem

s, limpid, volatile, inflammable, empyreumatic liquid, ich has received the name of acetone. This liquid may obtained, also, by dry distillation, from any dry acetate an alkali or alkaline earth; also by heating sugar of id with quicklime, by means of an iron bottle. When etone is distilled with half its volume of fuming sulphuric id, upon the liquid which passes into the receiver a yelv oil swims, which, after being washed with water, is :tified. The first portions contain a little acetone, which removed by redistillation, by means of a water bath. 3099. This oil is mesityl, being a colourless, oleaginous, lammable liquid, having a feeble odour, recalling that of rlic. It is lighter than water. With alkalies it underes no change. With sulphuric acid, nitric acid, and lorine, its habitudes resemble those of benzule. Its comsition is equivalent to two atoms of acetone, less two ms of water.

 $2C^3 H^3O = C^6 H^6 O^2$ Two atoms of acetone Deduct two atoms of water H<sub>3</sub> O<sub>3</sub> Co H And we have mesityl

4000. Acetone was inferred to be an hydrated oxide of esityl; but Dr. Kane, the author of the inference, has mitted that there are not sufficient grounds to justify n in retaining that idea of its composition. s peculiar and useful powers as a solvent. Many salts nich are soluble in both alcohol and water, are insoluble acetone, especially chloride of calcium and hydrate of tash. It burns with white flame, and has nearly the me density as alcohol. Its taste is disagreeable, having me analogy, however, with that of peppermint.

4001. Metascetone, C<sup>6</sup> H<sup>5</sup>O, is the name given to a coırless, volatile, fragrant, inflammable liquid, soluble in alhol and ether, but insoluble in water, and which boils at It may be considered as two atoms of acetone, nus one atom of water, C<sup>6</sup> H<sup>6</sup> O<sup>2</sup> — HO = C<sup>6</sup> H<sup>5</sup>O, me-

scetone.

4002. This liquid is generated simultaneously with acene, when one part of sugar, and eight parts of powdered ick-lime, are subjected to distillation.

ler sugar. But where a radical only furnishes the elementary ingredients to ther compound, or derives its ingredients from one, I do not conceive that any nexion in classification should be made between it and the substances whence it btained, or to the formation of which it contributes.

4003. Mesityl forms various compounds with the basecigen bodies, which it is not deemed proper to describe here. With sulphuric acid it forms a compound which affords soluble salts with baryta and lime.

Reflections on the Relation or Analogy between Acetyl, Ethyl,
Amide, and Ammonium.

4004. By the addition of an atom of water, HO, to ammonia, NH<sup>3</sup>, an oxide of ammonium is produced, NHO, which is the base of ammoniacal oxysalts (1116). In the manner it was supposed by Boullay and Dumas, that by the acquisition of an atom of water, etherine, a hydruret of carbon (1267), was enabled to play the part of a base a the neutralization of oxacids. This idea was, for some time, generally sanctioned, and hence, in the last edition of the Compendium, etherine was represented as the base of all the ethers which have, in this edition, been represented as having ethyl as their radical, and its oxide for their base.

4005. It has already been mentioned (1109), that agreeably to the doctrine advanced by Berzelius, and generally adopted, in the salts formed by presenting ammonia to be quid acids, the elements of the resulting base exist, not at a hydrate of ammonia, but as an oxide of ammonium. So far as an analogy with the habitudes of ammoniacal compounds would influence the view adopted, a corresponding conception would be created, that in etherial compounds the base should be an oxide of ethyl, not a hydrate of etherine. Besides the correspondence thus existing, there was no small analogy between the relation borne by amide to ammonium, and acetyl to ethyl: the only discordancy being, that the susceptibility of forming acids, displayed by acetyl, has not been observed in amide.

# Of the Compound Hypothetical Radical, Kacodyl, C' H'AS, Symbol Kd.

4007. The protoxide of kacodyl constitutes a fetid, viralently poisonous, etherial, spontaneously inflammable, vols-

tile, limpid liquid, long known as the fuming liquor of Cadet, its discoverer. This liquid, now called alcarsin, is obtained by distilling dry acetate of potash with an equal weight of arsenious acid. By digesting alcarsin, or oxide of kacolyl, in chlorohydric acid, chlorine taking the place of oxygen, a chloride of kacodyl results. From this the radical a separated, by reaction with metallic zinc, at the temperature of 230°, and removing the resulting chloride of zinc by water.

4008. Kacodyl is an etherial, limpid, spontaneously inlammable liquid having a nauseous odour. It sinks in water without dissolving, but is soluble in alcohol or ether. It boils at 338°. At a red-heat its vapour is resolvable into arsenic, olefiant gas, and light carburetted hydrogen.

4009. The following compounds are formed by this radical, of which it does not appear consistent to treat particularly here.

Kd O Alcarsin Oxide
Kd Cl Chlorarsin Chloride
Kd S Sulpharsin Sulphide
Kd Cy Cynarsin Cyanide
Kd O' + HO Alcargen Hydrated trioxide

4010. Agreeably to the preceding formulæ of the compounds of kacodyl, it may be seen that, excepting alcargen, they differ, in composition, only as respects one of their ingredients, a basacigen element, to the presence of which they owe the diversity of the names given in one of the lists.

4011. Alcargen, or kacodylic acid, differs from the rest

in holding an atom of water, HO.

4012. Liebig supposed the bodies in question each to consist of an atom of acetyl and an atom of arsenuretted hydrogen, As H<sup>3</sup>, not grouped into one radical; but Berzelius suggested that they were so grouped, and this Bunsen has proved to be true, by isolating kacodyl as above described.

4013. It may, however, be well to point out, that the composition of kacodyl is consistent with the idea of Liebig, since an atom of acetyl, - - C<sup>4</sup> H<sup>3</sup> and an atom of arsenuretted hydrogen, - As H<sup>3</sup> are equivalent to an atom of kacodyl, - C<sup>4</sup> H<sup>6</sup> As

1014. I object to the unmeaning names above given, as not conveying any idea of composition. Hence I shall

use those which indicate the composition.

4015. Alcargen, more significantly called kacodylic and, or hydrated trioxide, agreeably to the nomenclature which would make hydrous sulphuric acid a sulphate of water, would be a kacodylate of water.

### Of Methyl, C2 H3.

4016. After it had become evident that the etherial compounds, derived from the reaction of alcohol with acids or halogen bodies, had all a common compound radical, chemists were naturally led to infer, that there might be other series, similar in their nature, having the same electronegative ingredients united with other compound radicals. These speculative inferences first received a practical verification, from the labours of Dumas and Peligot respecting the composition and combinations of pyronic spirit, obtained from the products of the destructive distillation of wood or other organic products. From the seventigations of these chemists it has been made evident, that pyroxilic spirit is the alcohol of a series of compounds

having methyl as a radical.

4017. The compounds of methyl with the basaciges class, and those formed between its oxide and acids, are produced by reactions with methylic alcohol or ether, or their products, similar to those by which analogous compounds with ethyl are effected. There is, likewise, a great analogy in the properties of the two series; yet methylic ether (or in other words the oxide of methyl the compound which is the congener of ether proper), is gaseous, in lieu of existing like ether as a liquid. Moreover, a altrated oxide of methyl, or a true methylic nitric ether, is readily generated when wood spirit (hydrated oxide of methyl), is presented to nitric acid. This etherial compound has no congener among those of ethyl, because the reaction, between nitric acid and alcohol, is attended by a reciprocal decomposition, by which hyponitrous acid # evolved and combines, while nascent, with oxide of ethylexisting in an undecomposed portion of the alcohol-Hence it arises, that hypomitrous ether is generated ustead of nitrated oxide of ethyl. On the other hand so hyponitrite of the oxide of methyl, results from the reacion of nitric acid with wood spirit; since the decompoition, requisite to the development of hyponitrous acid, loes not ensue. Consequently, Liebig alleges that no congener of hyponitrous ether exists among the etherial combounds of methyl.

4018. I have recently been enabled to fill up this interral in the methyl series, by subjecting wood spirit to a hy-

ponitrite, in contact with a diluted acid.

# Of Formyl.

4019. Formyl has a relation to methyl, similar to that which acetyl has to ethyl. In either case, there is a radical differing from another, only by the subtraction of two

atoms of hydrogen.

4020. The hydrated oxide of formyl is inferred to exist in a liquid, obtained by the reaction of two parts of wood spirit with three of sulphuric acid, three of water, and two parts of peroxide of manganese. An inflammable, etherial, colourless liquid, of an agreeable aromatic odour and susceptible of solution in three parts of water, was thus procured. This liquid has been inferred to be a compound of two atoms of oxide of methyl, and one of hydrated oxide of formyl.

4021. There are in the formyl series no compounds corresponding to aldehyde, or acetous acid. The only exide is that long known as formic acid, from its having been first obtained from ants. This acid is obtained from formyl, as acetic acid from acetyl, by the addition of three

atoms of oxygen.

4022. Agreeably to Liebig, three chlorides of formyl have been isolated. The perchloride has been known for a good while, having been obtained by distilling alcohol with hypochlorite of lime. It was obtained about ten years since in this country by Gurthrie, and for some time confounded with the etherial oil of olefiant gas, which is now considered by Liebig as the chlorohydrate of the chloride of acetyl.

# Amyl, C10 H11.

4023. A peculiar liquid was noticed by Scheele to accompany potato spirit. Subsequently, by Pelletier, Cabours and D mas, it was inferred to be the hydrated oxide of a peculiar compound radical, to which the name at the

head of this article was given. It follows that this liquid must be a congener of alcohol, its formula being C<sup>\*\*</sup> H<sup>\*\*</sup> + HO.

4024. The amyl series of compounds corresponds with those of other radicals to a certain extent, but is upon the whole very incomplete, having no oxide to occupy the place of a congener of ether. Even the chloride does not appear to be permanent per se. The bromide and iodide are more enduring, and in their habitudes somewhat analogous to corresponding combinations in the series of other radicals.

4025. Yet in the case of sulphoamylic acid, the sulphose is well supported to other etherial double sulphose, such as sulphovinic acid, and there have been formed approamylates capable of decomposition and of reproducing

the hydrated oxide, potato spirit.

4026. An amylic acetic ether has been produced, by distilling two parts of acetate of potash, one part of potash spirit, and one part of sulphuric acid. As respects in flammability, volatility and insusceptibility of mixture with water, the amylo acetic ether is truly etherial in its nature.

4027. By the substitution of two atoms of oxygen for a like number of hydrogen, effected by treating potato spirit with hydrate of potash, a change in composition arises analogous to that by which alcohol is converted into acetic acid. An acid is in this way created, called valerianic, in consequence of its being identical in properties and composition with that extricated by distilling water from the root of valerian.

4028. This acid was produced, also, by causing potate spirit to fall slowly in successive drops upon platinum black duly heated. Peculiar liquids, somewhat etherial in their properties, have been evolved from potato spirit, of which the one C<sup>20</sup> H<sup>17</sup> Cl<sup>3</sup> O<sup>2</sup> seems to be a congener with chloral, the other with olefant gas the hydruret of carbon of Liebig.

# Glyceryl, C<sup>6</sup> H<sup>7</sup>.

4029. The wonderful fabric of scientific knowledge for which we are indebted to the skill, sagacity and ingenuity of modern chemists, is formed in part of materials which are altogether new, and in part of such, as although long known, owe nearly all their present theoretic value to the

hich they have latterly been made to answer in the fabric to which I have alluded.

O. In the preceding account of the amyl series it e noticed, that a liquid long since distinguished by e, and known under the name of oil of potato spirit of potatoes, has latterly been dignified with a place

the congeners of alcohol.

1. In glycerine, C<sup>6</sup> H<sup>7</sup> O<sup>5</sup> + HO, the hydrated oxide compound radical glyceryl, we find, in like manner, bound of similar antiquity, and, as respects its disr, of like origin; having been well known since the f Scheele, as the sweet principle of oils. For the hich it now occupies, the scientific world is indebted vreul and Pelouze.

2. Anterior to the labours of Chevreul, an erroneous existed that the process of saponification consisted ing more than a union between the alkali and oil; it was deemed to be a case simply of combination. istence in every oil of an electro-negative, and an -positive ingredient, the one performing the part of

, the other of an acid, was not imagined.

3. The oxide of glyceryl is the base common to a ty of vegetable and animal fixed oils, whether liquid solid state, denominated fat, being liberated during iling of those substances with fixed alkalies, as in cess of saponification. It is best prepared by sapooil of olives with litharge, separating the resulting n of glycerine, and precipitating any dissolved lead phydric acid (897, 899).

1. Glycerine is said to be deficient of two properties ing to its alcoholic congeners, solubility in ether, sceptibility of distillation without decomposition. It et, colourless, and inflammable; of the density of

being about one-fourth heavier than water.

i. It does not appear that there are any other imt compounds formed with this radical by the basacidies or the acids, so as to be productive of compound congeneric with those so formed by most of the therefiable compound radicals. There is, neverthecongener of sulphovinic acid in sulphoglyceric acid, properly called the double sulphate of the oxides of yl, and of hydrogen.

# Cetyl, C38 H33.

4036. Of cetyl it may be sufficient to say, that it is perfectly analogous as respects the part which it performs a spermaceti, with that performed by glyceryl, as the resistance.

cal in the base of the fixed oils generally.

4037. The diversity of such oils, in other cases, is produced by variation in the acids with which the cride of glyceryl in them severally is combined. Spermaceti has been represented as a solitary instance in which a charge of properties results in a concrete fixed oil, from a prolibility in the hydrated oxide constituting the base, while the acids, combined with this base, are those which have been described as entering into the composition of oleginous products in general. Recently, this view of the subject has been controverted by Smith. Silliman's Journal, October, 1842. (See 5055, page 426.)

4038. The hydrated oxide of cetyl, C H O + HO, my be elaborated from spermaceti by saponification, in a make resembling that by which glycerine is obtained. It has been designated by the name ethal, a word made up of the initials of alcohol and ether. It differs from other alcoholic hydrated oxides, in being deficient of that solubility in water which is one of the most striking and distinguishing attributes of alcohol proper. It differs also in being solid until heated to 118°. The analogy with glycerine fails as respects taste, being insipid; also in this, that gly-

cerine is soluble in water, and insoluble in ether.

4039. Cetyl has not been isolated; but by repeated distillation with anhydrous phosphoric acid, ethal has been made to yield an inflammable liquid compound, C<sup>2</sup> H<sup>2</sup>, having to it a relation analogous to that which etherine or etherole, C<sup>4</sup> H<sup>4</sup>, has to ethyl, C<sup>4</sup> H<sup>5</sup>. Cetene, as this liquid is called, seems to be of the nature of an essential oil, since it may be distilled. It requires, however, the high temperature of 527° for this purpose.

4040. Cetyl coincides in habitudes with the other compound radicals of this class, as respects the formation of double sulphates, analogous to the sulphovinates. It also forms a chloride capable of being distilled, and by the substitution of three atoms of oxygen for two of hydrogen, is converted into an acid, denominated ethalic, C<sup>20</sup> H<sup>20</sup> O,

which is a congener of acetic acid.

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# OF NUTRITIOUS VEGETABLE SUBSTANCES DEVOID OF NITROGEN.

4041. Under this head I place gum, sugar, fecula, and ignin. Immediately, this last mentioned substance is rather food for worms than for man; but it will be seen that lig-

nin may be converted into sugar.

4042. The substances above enumerated might be treated as hydrates of carbon, agreeably to the suggestion of Prout (2096), were it not that their properties do not warrant the idea, that the hydrogen and oxygen are more intimately allied to each other, than to the carbon.

# Of Gum.

4043. Substances known under the generic name at the head of this article have certain properties in common, but vary with the tree by which they are generated. Some, like gum arabic, or gum senegal, are perfectly soluble in water; while others, like tragacanth, are capable only of forming a paste with the same liquid. Those of the first mentioned kind are susceptible of rapid desiccation and induration, by access of atmospheric air, while the others give up water, comparatively, with reluctance. They are all distinguished from resins, which they resemble externally, in being insoluble in alcohol, ether, or essential oils. They differ from sugar in the want of sweetness, and from starch in not being coagulable by heat.

4044. Guerin, in an elaborate treatise on gums, divides them into three classes:—1. Arabin, of which gum arabic is the type, soluble in cold water. 2. Bassorin, which swells into a jelly, but does not dissolve in water: gum bassora, or tragacanth, may exemplify this class. 3. Cerasin, from the gum of the cherry-tree. Cerasin is also insoluble in cold, but soluble in boiling water, and when treated with nitric acid, gives about one-fourth less mucic

acid than bassorin.

4045. Of arabin, by his analysis, the formula is C<sup>6</sup> H<sup>5</sup> O<sup>5</sup>. Gum senegal, and the soluble parts of gum tragacanth and bassora gum, consist of arabin.

4046. Of bassorin the formula is C<sup>10</sup> H<sup>11</sup> O<sup>11</sup>.

4047. Cerasin appears to be metamorphic arabin; for it has precisely the same composition, and is changed into it by solution in boiling water. The gums of the cherry, apricot, prune, peach, and almond tree, are of this kind.

4048. Berzelius employs the word mucilage to designate that species of matter which is exemplified by the bassorin of Guerin. Varieties of this kind of gum seen in infusions of flaxseed, of slippery elm, and pith of sassafras. This use of these terms is not adopted by Tuner, Kane, or Graham. The principal difference between gum and mucilage, agreeably to general acceptation, seems to be, that mucilage is not susceptible of spontaneous her-dening by desiccation. Graham admits only of two general of gums, exemplified by gum arabic and gum tragacestic By Kane, they are treated of under three heads—archin, cerasin, and dextrine, or artificial gum. This last metioned variety is obtained from starch, and does not w pear to have higher pretensions to be ranked as a gum than the modification of starch by heat, known as British gum. Substances which come under the name of gum, agree in general properties; yet there are scarcely at two which are quite similar. Gum arabic is deemed to be the most perfect specimen of the substances bearing this name.

4049. Berzelius considers the reaction of a solution of this substance with a solution of the silicate of potash, as the most striking characteristic of its properties. One portion of it forms, with one part of the alkali and all the acid, a triple compound, which precipitates; while another portion of the gum, and the remainder of the potash, combine and remain in solution.

4050. Gum arabic differs from other gums in combining with the sesquioxide of iron, and forming a compound insoluble in water, but soluble in acids. A solution of gum arabic in 1000 parts of water, being mixed with a solution of the sesquioxide of iron, yields, in 24 hours, a yellow precipitate. This species of gum also combines with, and precipitates the protoxide of mercury from the nitrate. There appears to be no important difference between gum senegal and gum arabic.

# Of Sugars.\*

4051. Under this head I would place two genera of substances; crystallizable sugars, and syrups incapable of

<sup>&</sup>quot;Liebig treats of sugars under the general head of an "appendix to the combinations of ethyl and acetyl." His commentator, Gregory, alleges that they are "thus treated of, since from them are derived all the compounds of ethyl; and, also, be-

crystallization, and which might be called suavin. Of the former, sugar candy, and the crystals found in raisins and honey, are specimens. The latter are exemplified by the uncrystallizable syrups of raisins and of honey; also the sweet matter of the sweet potato, and the uncrystallizable syrup of the sugar cane, known as molasses.

4052. The qualities, both of crystallizable and uncrystallizable sugars, vary with the plants from which they are produced. In the power of imparting sweetness to infusions, the crystallizable sugar of the cane is pre-emi-

Bent.

4053. As sugar has been found to be very susceptible of yielding alcohol by fermentation, this property has been made the basis of defining the meaning of the word, so that every substance capable of the process alluded to, is to be considered as sugar, whatever may be its taste, or however it may differ in its properties from the substances assually called by the name.

4054. Thus the fermentable "wort" of distillers or brewers, the uncrystallizable juices of fruits, a substance found in mushrooms or ergot, also an insipid matter found by Thenard in diabetic urine, are all to be considered as consisting of sugar, so far as they are capable of yielding

alcohol by fermentation.

4055. I am reluctant to employ words in a sense different from that in which they are generally understood. Agreeably to usual acceptation, sweetness is an indispensable attribute of sugar. Sugary and sweet are synonymous. "As sweet as sugar" has long been an expression conveying the idea of superlative sweetness.

4056. Chemists have erred, I think, in assuming that

nates the uncertainty in which we are as to their true constitution, renders it im-

possible to arrange them on scientific principles."

That ethyl compounds are derived from sugars, might be a reason for treating of them under sugars; but I cannot perceive the converse to be true. But as aldeby-bus or acetons acid, and acetic acid, are placed under the head of acetyl, and the tempounds of mesityl are derived from acetic acid, a compound not necessarily derived from sugar, if the reason above given were sufficient for placing sugars under ethyl, it is, on that same ground, improper to place them under acetyl, since this radical is not necessarily a product of sugar.

In reply to the last sentence quoted, it might be demanded, why inability to arrange segers upon scientific principles, justifies their being placed under the head selected, in preference to any other: whether every set of substances which cannot be arranged on scientific principles, are to be discussed under the joint head of com-

binations of ethyl and acetyl?

The best justification which occurs to me for any connexion between cane sugar and sectyl is, that when anhydrous it is isomeric with acetyl, one atom containing three of this radical, acetyl, Ca Ha + 3 ms CB Ha Oo.

nothing besides sugar is susceptible of the vinous fermentation. The conversion into alcohol of the insipid product of diabetes, which has been treated as sugar, because proved to be susceptible of the process in question, might with more propriety, as I conceive, be deemed to demonstrate that this process may be undergone by substances which are not sufficiently of a saccharine nature to merit

the name of sugar.

subjected to a ferment, at a certain time before its conversion into alcohol, it affects polarized light in the same way as grape sugar. Hence it is inferred, that cane sugar is not directly susceptible of the vinous fermentation; and that of all sugars, that of the grape only is capable of immediately undergoing that process. It follows, that if the contested definition be not disregarded, the sweet crystallizable matter extracted from the cane, hitherto considered as the most perfect of the sugars, must be deprived of its title, and occupy a place on a level with starch, as being, like this substance, incapable of the vinous fermentation, without a previous transformation into grape sugar.

4058. Liebig enumerates the following varieties of sugar. Cane sugar, grape sugar, lactin or sugar of milk, uncrystallizable sugar, and sugar of mushrooms. To these Graham adds, insipid diabetic sugar, manna sugar or man-

nite, and liquorice.

4059. As a good account of the sources of the sugar of commerce, and the means by which it is elaborated may be found in the United States' Dispensatory, it will be doubly inexpedient to extend in this treatise the information given, beyond its chemical composition and habitudes.

4060. Of cane sugar, perfect specimens are seen in the best double refined sugar, and in colourless sugar cardy. Its specific gravity is 1.6. At 350° it fuses into the well known form of barley sugar, which, by exposure to air, is

alleged to become white, opaque, and crystalline.

4061. Exposed to the temperature of 650°, by losing as atom of water besides that of crystallization, sugar is transformed into the dark brown substance called caramel. Thus obtained, caramel is not entirely exempt from undecomposed sugar and other impurities, but may be freed from them by solution in water, and precipitation by alco-

. The precipitate thus created, when dried, forms a ck or dark brown powder, which may be redissolved in It is insipid, not fermentable, and neither acid nor aline. Caramel is used to deepen the colour of fernted or spirituous liquors. During its decomposition heat, fumes are emitted by sugar, which not only disse, but, as I believe, neutralize fetid emanations. For solution, cane sugar requires one-third of its weight of d water, but the effect of this liquid at a high temperae, is rather that of lowering the point of fusion, than ing as a solvent; since, at the temperature of 350°, sur liquefies per se, and of course may liquefy with the mitest proportion of water which can be added. liquefaction is due to heat rather than to water.

4062. If a concentrated solution of sugar be subjected, some time, to the temperature requisite to vaporize the ess of water, under the whole pressure of the atmosre, it is changed by degrees into uncrystallizable sugar. nce, of late years, the vaporization is aided by a reduca of atmospheric pressure, by means of an air pump.

**72.**)

1063. Sugar combines with some salts; and acts feebly an acid, so far as to unite with some bases. oluble compound, formed by anhydrous sugar with oxied lead, the base is a dioxide. With one atom of baa, sugar forms a crystalline compound; with common

t it forms crystals readily soluble in water.

1064. Berzelius alleges that an aqueous solution of sudissolves the carbonate and subacetate of copper, ing rise to a green liquid, from which the metal is preitated by sulphydric acid, or cyanoferrite of potassium, not by alkalies. When boiled with solutions of cuous salts, it causes the reduction of the copper.

1065. Several products are obtained by the reaction of ious acids, either dilute or concentrated, with the various ds of sugar; also by their reaction with alkalies. These ducts being complicated in their nature and of little

ctical utility, I shall not treat of them here.

Grape Sugar.
Crystallized, C<sup>12</sup> H<sup>14</sup> O<sup>14</sup>; Anhydrous, C <sup>12</sup> H<sup>12</sup> O<sup>12</sup>.

1066. Crystals of this sugar may be seen in raisins, in at are called candied sweetmeats, and in honey, in either 52

of which it forms the least fluid portion. Fruits generally owe their sweetness to its presence. The sugar formed from starch by digestion with diastase, or sulphuric acid,

is of this species; and also the sugar of diabetes.\*

4067. Grape sugar may be obtained in crystals from grape juice, by neutralization with chalk, clarifying with albumen, evaporation, and subsequent repose: also from diabetic urine, by evaporation to dryness, by means of a water bath, washing the resulting crystalline mass on a filter with cold alcohol until it becomes white, and repeated re-solution and recrystallization.

4068. It is remarkable, that notwithstanding the amlogy between cane and grape sugar, they differ much in their chemical qualities, as shown by their habitudes with chemical reagents. Strong mineral acids, which react but feebly with grape sugar, readily decompose cane sagar. With alkalies an opposite result ensues. The compounds which are formed by these sugars respectively, with bases, are quite different.

4069. From an alcoholic solution, grape sugar crystallizes in transparent square tables or cubes; from an aqueous solution, it consolidates into a spongy mass of crys-

talline grains.†

1 See fermentation; also (4071).

# Sugar of Milk, or Lactin.

4070. The formula of crystallized sugar of milk is C<sup>24</sup> H<sup>24</sup> O<sup>24</sup>, or C<sup>24</sup> H<sup>19</sup> O<sup>19</sup> + 5HO. By a heat of 248° it loses two atoms of water, and by 302°, five atoms. (Berzelius.) It is obtained by evaporating the whey of milk, and purifying the first crystalline product by animal charcoal and recrystallization. It forms white, semi-transparent, quadrangular prisms, which have the density 1.543. They are soluble in five or six parts of cold water, and in two and a half parts of boiling water. The taste of the crystals is very feeble, being inferior, in sweetness, to that of their solution. Sugar of milk is unalterable in the air, or by a heat under 212°, and is insoluble in alcohol or ether. When milk is exposed to a temperature of from 95° to

<sup>\*</sup> Dumas has proposed that grape sugar be called glucose; but as Liebig alleges that all sugars, even that of the cane, have to be converted into grape sugar in order to be rendered susceptible of the vinous fermentation; it would seem, consistently with the received definition of sugar (4053), as if cane sugar should be called giscose, yielding the name of sugar to the sweet matter of the grape.

it undergoes the vinous fermentation, generating all, while its sugar disappears. But it is presumed that atter is converted first into grape sugar, probably the influence of the free acid, which, being formed, es the milk. Milk sugar forms two compounds with of lead, of which the formulæ are C<sup>24</sup> H<sup>19</sup> O<sup>19</sup> + 5PbO, C<sup>24</sup> H<sup>19</sup> O<sup>19</sup> + 10PbO. (Berzelius.)

# Mushroom Sugar.

71. This sugar, of which the formula is C<sup>12</sup> H<sup>13</sup> O<sup>13</sup>, ding to an analysis by MM. Liebig and Pelouze, obtained by M. Wiggers, by subjecting the tincture ergot of rye to water. It is crystallizable and solutwater and alcohol, but insoluble in ether. Mush-sugar is also fermentable by yeast, and diffuses the rof caramel when carbonized by a high temperature. only property by which this sugar is distinguished the ordinary species is, that it does not throw down xide of copper from a boiling solution of the acetate.

e Fermentable Matter of Diabetes, called Insipid Sugar.

72. It has been stated (4052), that a substance was ned, by Thenard, from the urine of diabetes insipidus, subsequently, by Bouchardat, from the same source, I was insipid, or only faintly sweet. The aqueous on of this sugar was fermentable by yeast, and susple of being converted into the sugar of grapes by dialphuric acid.

## Liquorice Sugar.

73. The inspissated juice of the root of the Glycyr-glabra contains a species of unfermentable sugar, 1 may be obtained by clarifying the juice with albuprecipitating the sugar with sulphuric acid, washing recipitate with water, dissolving it in alcohol, which ates some undissolved albumen, and then decompt the sulphate of liquorice sugar by carbonate of pot-After evaporation, the sugar remains as a yellow lucent mass, cracked in all directions, and easily ded from the vessel in which it has been desiccated. Frice sugar is capable of forming soluble or sparingly le compounds, with both the mineral and vegetable. It also combines with bases.

## Manna Sugar, or Mannile.

4074. The formula of manna sugar is C<sup>6</sup> H<sup>7</sup> O<sup>6</sup>, according to the analysis of Oppermann and of Liebig. Manua is in oblong globules or masses, of a yellowish-white colour, being an exudation from various trees, principally the fraxinus ornus, and encalyptus mannifera of New South It exists also in the juices exuded by cherry and plum trees, in those of various kinds of mushrooms, and of celery and other roots. Manna sugar may be prepared by dissolving the manna of the shops in boiling alcohol, and allowing the solution to cool. It may be purified by repeated crystallizations. Mannite crystallizes in slender, colourless, four-sided prisms, of an oily lustres It has a slightly sweet taste, forms, with water, a solution which is not fermentable. It is anhydrous, and may consequently be heated to redness, without any loss of weight. Its aqueous solution dissolves oxide of lead. Nitric acid coverts mannite either into oxalic, or saccharic acid; but not into mucic acid. Mannite is also one of the products of the vinous fermentation of cane, or grape sugar.

## Fecula, or Starch.

4075. A substance, of which starch is a good specimen, and of which the generic name is fecula, may be obtained from the meal or flour of grain, and from the tubers of the potato, and various other vegetables. It is found in commerce under the names of sago, tapioca, arrow-root, &c. Of the sources of these varieties of fecula, an excellent account is given in the United States Dispensatory. It is more or less a constituent of vegetables in general. When the farinaceous matter, procured from such sources by rasping or grinding, is washed, the fecula is suspended, and subsequently deposited. Where there is vegeto-animal matter, as in wheat flour, fermentation is employed to get rid of this substance.

4076. It was discovered by Leeuwenhoeck, with the aid of a microscope, in 1716, that starch consists of globular grains, each enveloped in a tegument, pocket, or sac, differing from the internal mass. In 1825, these observations were confirmed and extended by Raspail, who also

observed that the envelope, or tegument, was insoluble in water, while the interior portion was soluble in this liquid. Agreeably to the microscopic observations of this last mentioned author, the sizes of the globules of fecula vary with the plant whence it may be derived. Those of the potato did not exceed in diameter 110 of an inch; those of wheat 110 of an inch; and of arrow-root 110. As, according to Payen and Persoz, the tegument does not form more than four or five thousandths of the weight, the internal portion may be considered as characterizing the whole, uninfluenced to any important extent by the tegumentary matter.

4077. Fecula is blackened by a certain quantity of iodine, becomes blue with less, and violet with still less. The iodide of starch becomes colourless at a temperature less than 200°, and if not made to reach the boiling point, regains its colour on cooling.

4078. Starch does not combine with cold water, but forms a viscid solution with hot water. It is neither dis-

solved nor acted upon by alcohol or ether.

4079. Fecula dissolves in nitric acid without heat, and when heated with it is converted into oxalic acid. A slight torrefaction changes its nature, so that it may be used as a substitute for gum. Triturated with potash, fecula acquires the property of dissolving in cold water. The solution is clouded by acids.

4080. Its solution in hot water is precipitated by subsalts of lead, and in cold water by an infusion of galls.

## OF DIASTASE,

And of the Conversion of Fecula into Dextrine and Grape Sugar.

4081. Boiled in water, constantly replenished for nearly forty hours, with between \( \frac{1}{10} \) and \( \frac{1}{10} \) of its weight of sulphuric acid, fecula is converted into grape sugar. A similar change is alleged to have ensued partially in starch, which was made into a paste with twelve times its weight of boiling water, and kept for two years. By the addition of the glutinous matter obtained by washing wheat dough, and the application of a heat between 122° and 167° Fah., a similar result is said to have been attained in about ten or twelve hours.

4082. It is well known to those who are acquainted with the manufacture of whiskey from grain, that a portion of malt is necessary to render the wash or wort susceptible of the vinous fermentation; and that the product is much affected by the circumstances under which the infusion of the grain is accomplished. Nearly thirty years ago, my late friend, Col. Anderson, who had distinguished himself by his ingenuity and sagacity in improving the processes and apparatus of our American distilleries, expressed to me an opinion, that the mixture of farina and water became sweeter towards the close of the process of infusion, and that he believed a chemical change was induced, by which more or less sugar was generated. The inference of our ingenious countryman has been fully justified by the researches of Payen and Persoz, whence it appears that, by digestion with malt, fecula is at first partially changed into a sweetish gummy matter, called dextrine, and that this matter is afterwards converted into grape sugar. Destrine has received its name from a peculiar influence which it exercises upon the plane of polarization, during the passage of light.\* It may be considered as holding, as respects its properties, an intermediate position between fecula and grape sugar.

4083. The sugar-producing property thus existing in malt, has been traced to a peculiar principle called diagrass, which exists therein in a proportion not exceeding a five-hundredth. It is obtained by moistening ground malt with half its weight of water, and exposing the mass to pressure. The exuding liquor is mingled with a quantity of alcohol of 840°, by which the diastase is thrown down impure. By three successive solutions in water, and precipitations by the same means, with subsequent exposure on a glass pane, in thin layers, to a current of air about

When light, polarized by reflection from the surface of a plate of black glass, of from the surface of a pile of superposed plates of transparent glass, reaches the systhrough a disc of tourmain, a solution of dextrip being interposed in a tube between the reflecting plate and tourmain, the light does not disappear in those positions of the tourmain in which light would be completely absorbed without the interposition of the solution of dextrine, but primatic colours are produced which follow a certain order, if the plane of polarization is turned from left to right. It is by the order of these colours, that a liquid is said to polarize light to the right or to the left. The solution of starch polarizes to the right, and that of dextrine considerably more as the same direction, while a solution of once sugar produces the succession of colours in an inverse order, and is said therefore to polarize to the left. The program of changes may thus often be observed in a solution of starch, the justice of plants, and other organic fluids.—Graham, 743.

a white amorphous solid matter. Diastase does not er gum, sugar, gluten, nor albumen, nor the teguments fecula, but operates surprisingly, as above described, on ula proper. This change is effected without any abption of the air, or any evolution of gaseous matter. It y take place either in pleno or vacuo. An infusion of parts of starch in 39 parts of water, at about 90° Fah., ng mixed with 6.13 parts of diastase, dissolved in 40 ts of cold water, and digested afterwards for an hour, a temperature between 90° and 100°, gave 86.91 parts sugar. At the temperature of 158°, one part of diase will convert 2000 parts of starch into sugar.\*

1084. When sulphuric acid is employed in lieu of diase, if, by confinement, the temperature and pressure are sed (192), less sulphuric acid will suffice. Less time is uisite when care is taken to prevent too rapid refrige-

on.

1085. If a paste, made by subjecting starch and water sbullition, be gently poured into a boiling dilute solution sulphuric acid, the pasty consistency soon disappears. like manner, starch paste loses its gelatinous character en mingled with malt wort, and if kept at a temperature ween 190° and 200°, becomes, at the end of some hours, verted into grape sugar.

086. In proportion as the diastase saccharifies the ch, it disappears itself; and when the solution no ger acts on a fresh portion of starch, the presence of stase cannot be detected in it. The reaction is probachemico-electric, and if understood, would throw light

a multitude of phenomena.

t Gregory's Turner, 943. Graham, 744.

ostance, soluble in water and in weak alcohol, but inible in absolute alcohol. It is not known to enter into
ibination with any substance.† It received its name
in sorthul, I separate, in reference to separation of the
elope of the starch globules (4065).

# Lignin.

1088. The tasteless, inodorous, insoluble, but tenacious es of wood, hemp, cotton, or flax, and other plants,

<sup>•</sup> Graham, 745. Annales de Chimie et de Physique, Vol. 53, p. 73.

have been deemed to consist of a peculiar vegetable substance, called lignin, from lignum, the latin for wood. The formula of lignin, dried between 300° and 350°, is C<sup>12</sup> H<sup>1</sup>O<sup>2</sup>

(Prout).

4089. Graham alleges, that it constitutes about 95 per cent. of baked wood, and that it may be obtained in purity by treating the sawings of wood, paper, or the fibre of lint, cotton, hemp, &c., successively with ether, alcohol, water, diluted acid, and a caustic alkaline solution, so as to dissolve and remove all the matter soluble in those mensure. Wood consists of an association of capillary tubes, in which, after it is desiccated, agreeably to the observations of Hartig, a quantity of starch remains, in spherical grains of a grey colour. Hence by triturating it, in the state of fine saw-dust, with water, from one-fourth to one-fifth of its weight of starch may be obtained.

4090. If Payen is to be credited, wood consists of two organic principles, one of which is isomeric with starts, having the same formula, C<sup>12</sup> H<sup>10</sup> O<sup>10</sup>, being named cellulose by him. The other principle, which forms the tubes, is considered by the same author as the true lignin. Cellulose was obtained by subjecting sawings of beech wood to several times its weight of the most concentrated nitric acid, which leaves that principle, while it dissolves the lignin. Cellulose is dissolved by concentrated sulphuric acid without blackening, and is then converted into dextrine. The formula of lint, hemp, straw, and linen cloth, was found by Payen to be C<sup>35</sup> H<sup>24</sup> O<sup>20</sup>. Oak wood, by the ana-

lysis of Gay-Lussac and Thenard, is C36 H22 O22.

4091. When hemp, straw, &c., are added cautiously to concentrated sulphuric acid, so as to prevent elevation of temperature, not only is dextrine created, but also lignosulphuric acid, analogous to benzo-sulphuric acid, which forms a soluble salt with baryta, or with oxide of lead.

4092. The dextrine formed when lignin is dissolved in sulphuric acid, is converted, by dilution and boiling, into

starch sugar.

4093. Saw-dust, gum, and starch, dissolve in the most highly concentrated nitric acid, without decomposing the acid; and, if immediately diluted with water, give a white pulverulent neutral substance, insoluble in water, which contains the elements of nitric acid, and is highly combustible.

34. To obtain grape sugar from lignin, twelve parts reds of paper or linen, or of wood shavings, are intividually incorporated by trituration with seventeen parts neentrated sulphuric acid (according to Vogel five), and one of water, carefully preventing any rise of trature. After twenty-four hours, the resulting tarry is to be dissolved in water, boiled for ten hours, neud with chalk, and being filtered and evaporated to a y consistence, the residue is to be left to crystallize. It is according to Brunner, 100 parts of fecula yield for crystallized grape sugar; according to De Saussure,

Agreeably to calculation, 100 of fecula, with four of water, should be productive of 120 of sugar. 100 of linen shreds produce 114 of sugar, according to

onot; or, according to Guerin, 115 parts.

16. It is worthy of remark, that the formula of crysed grape sugar may be made by adding to the forof lignin six atoms of water; to that of starch, four
i; to that of cane sugar, three atoms; and to that of
of milk, two atoms.

| a of lignin,<br>ms of water,      | C <sub>18</sub> H <sub>8</sub> O <sub>8</sub>   | Starch,<br>Four atoms of water,       |     |                        | O <sup>10</sup> |
|-----------------------------------|-------------------------------------------------|---------------------------------------|-----|------------------------|-----------------|
| lized grape sugar,                | C19 H14 O14                                     | Grape sugar,                          | C19 | H14                    | O14             |
| lized cane sugar, stoms of water, | C <sub>18</sub> H <sub>11</sub> O <sub>11</sub> | Sugar of milk,<br>Two atoms of water, |     | H <sup>19</sup><br>H • | O a             |
| ngar,                             | C19 H14 O14                                     | Grape sugar,                          | C14 | H14                    | O14             |

#### OF VEGETO-ANIMAL SUBSTANCES.

- this Head are included Gluten, Vegetable Albumen, getable Fibrin, and Legumen, or Vegetable Caseine.
- 7. Plants contain substances which have been desigas vegeto-animal, on account of their analogy with hite of egg, and the fibrin of animal matter. Nitroalways an ultimate element in them, and occasionalphur and phosphorus. As they are to be found in getables, to a greater or less extent, it appears proarrange them under the head of the general princifivegetables.

18. It had long been known that wheat dough, by enclosed and kneaded within a porous bag, while 53

subjected to water, might be resolved into a portion which would be washed away by the water, and an adhesive por-

tion left within the bag.

4099. Beccaria first drew the attention of chemists to the substance thus obtained. Subsequently, Rouelle, Jr., demonstrated the existence in the expressed juices of many plants, of a substance coagulable by heat, like animal albumen. This coagulable matter was, by Fourcroy, demod to be of the same nature as the albumen of eggs. Subsequently, Einhof demonstrated the existence, in rye, batter, peas, and beans, of two vegeto-animal substances; one resembling white of egg, the other, which he designated at gluten, was not considered as resembling any animal substance.

5000. It may be inferred, from the account of gluten given by Berzelius, that both Einhof and Tuddei subjected the gluten of Beccaria to boiling alcohol, and thus resolved it into two substances; one similar to albumen in its properties, the other soluble in alcohol, especially when boiling, and possessing, in a high degree, the adhesiveness and other properties by which gluten had been distinguished.

by Taddei, designated as gliadine, from yam, glue, the portion remaining undissolved, zimome, from gran, leaves. Berzelius treats the matter, soluble in alcohol, as gluten nearly pure, and the residue as vegetable albumen, and gives the following account of the sources and properties of gluten and the vegetable albumen with which it is associated.

#### Gluten.

5002. It owes its name to the adhesive property which it possesses, and which it communicates to wheat dough. It exists in the seed of the grape, and of the cerealia especially; also in those of leguminous plants, such as pear and beans, in which it is found in combination with starch and vegetable albumen. Its distinguishing characteristics are as follows. When isolated, it is almost insoluble in water. It is gluey when moist, yellow and translucent when dry. Ordinarily, it has an acid reaction with litmus, in consequence of the presence of acetic and phosphoric acid. It is soluble in alcohol, especially when boiling, and likewise in diluted aqueous solutions of acids, caustic alka-

eys, &c. It is precipitated from the latter by ferroiate of potash. With nut-galls it gives a precipitate, h is not redissolved even by ebullition.

## Vegetable Albumen.

03. It is found in the above mentioned seeds in comion with gluten; in seeds which yield emulsions, as, istance, in almonds; and likewise in the seeds of the is, where it is found in combination with an oil. It is in all vegetable juices which coagulate with heat. It table albumen is soluble in water, until coagulated by but is not soluble in alcohol; it is not adhesive, and esiccation becomes opaque, and of either a white, brown, or black colour. It dissolves readily in it is alkaline solutions, neutralizing their caustic taste, is precipitated by a great excess of acid. The prette is a chemical compound of albumen with the acid, le in water when pure, but less so when this liquid is ilated.

04. The aqueous solution of vegetable albumen is pitated by acids, by ferro-prussiate of potash, by chlo-of mercury, and infusion of galls; being, in these res, perfectly analogous to animal albumen.

O5. Gluten and vegetable albumen spontaneously uno decomposition, accompanied by an evolution of ama, a production of the acetate of ammonia, and likethe fetor which distinguishes the putrefaction of anisubstances. At a certain period of putrefaction, they
, whether separate or mixed, the smell of old cheese.

# Of the Gluten and Albumen of Wheat.

O6. If we make a thick paste of wheat with water, in rous bag, and knead this paste within the bag, under r, until this liquid is no longer rendered milky, there ins, finally, a gray coherent elastic residue. This lue consists mainly of a mixture of gluten and vegeta-lbumen, not quite free from other matter derived from wheat, and more or less of starch, which it is difficult move completely. This residue does not contain the le of the vegeto-animal matter of the wheat, a part g carried away by the water during the kneading of paste.

5007. To separate from each other the albumen and gluten proper, contained in the gluten of Beccaria, it is more cessary to subject it to boiling alcohol, till this liquid, on being filtered, is not made turbid by cooling. The alcohol dissolves the gluten proper, as well as another substance imperfectly known, leaving the vegetable albumen. The gluten is obtained by mixing the alcoholic solution with the water, and removing the alcohol by distillation. A liquid remains, in which the gluten floats in coherent voluminous flocks. A very small portion of gluten remains in

solution, combined with gum.

5008. The gluten being separated from the liquid, is of a pale yellow, and readily becomes agglutinated into mass, which sticks to the fingers, is clastic, insipid, and endowed with a peculiar odour. In dry air it becomes spontaneously polished on the outside, and of a deeper yellow, drying, by little and little, into a translucent mass of a very deep yellow, resembling dried animal matter-Alcohol dissolves the gluten, and the solution, which is of a pale yellow, being evaporated, the gluten remains in the form of a yellow transparent varnish. If the gluten be macerated in cold alcohol, it is whitened, and forms a milky solution, from which an insoluble matter is deposited. This is not gluten, though of a kindred nature, being soluble by the aid of ebullition; the resulting solution, when concentrated, acquires a mucilaginous consistence on cooling. Gluten dissolves in boiling officinal alcohol, and precipitates by refrigeration, without having lost its gluey quality. It is insoluble in ether, or in fixed out or volatile oils. If we subject it to acetic acid, it becomes, in consistency, mucilaginous, semi-liquid, losing its yellow colour. Mixed in this state with water, it gives a mucilaginous flocky residuum and a milky solution.

5009. From the investigations of Einhof, as stated by Berzelius, it appears that a matter, analogous to that above described as true gluten, may be obtained from rye, barley, oats, or even from maize, which, from the absence of any cohesiveness in its moistened meal, would not be supposed to contain any matter deserving to be distin-

guished as gluten.

5010. It will be perceived, from the preceding history of the opinions and observations of chemists, respecting the vegeto-animal matter obtained from wheat and the seeds

other vegetables, that the idea lately put forth by Lie, respecting the identity of their composition with anilabumen, has long been entertained, though never beer presented so forcibly to popular attention.

ion. Respecting the matter treated as gluten by Berius, Liebig advances views which are in some respects v, and somewhat discordant. I will here quote the lange of the author last mentioned:—

These nitrogenized forms of nutriment in the vegetable kingdom may be red to three substances, which are easily distinguished by their external charac-

Two of them are soluble in water, the third is insoluble.

When the newly-expressed juices of vegetables are allowed to stand, a separatakes place in a few minutes. A gelatinous precipitate, commonly of a green, is deposited, and this, when acted on by liquids which remove the colouring er, leaves a grayish white substance, well known to druggists as the deposite vegetable juices. This is one of the nitrogenized compounds which serves for autrition of animals, and has been named vegetable fibria. The juice of grapes pecially rich in this constituent, but it is most abundant in the seeds of wheat, of the cerealia generally. It may be obtained from wheat flour by a mechanical ation, and in a state of tolerable purity; it is then called glutes, but the glusproperty belongs, not to vegetable fibrin, but to a foreign substance present in quantity, which is not found in the other cerealia.

The method by which it is obtained, sufficiently proves that it is insoluble in ;; although we cannot doubt that it was originally dissolved in the vegetable

, from which it afterwards separated, exactly as fibrin does from blood.

he second nitrogenized compound remains dissolved in the juice after the sepan of the fibrin. It does not separate from the juice at the ordinary temperature, a instantly coagulated, when the liquid containing it is heated to the boiling

When the clarified juice of nutritious vegetables, such as cauliflower, asparagus, yel wurtzel, or turnips, is made to boil, a coagulum is formed, which it is absorptionable to distinguish from the substance which separates as coagulum, the serum of blood or the white of an egg, diluted with water, are heated to oiling point. This is regetable albumen. It is found in the greatest abundance rtain seeds, in nuts, almonds, and others, in which the starch of the graminese placed by oil.

The third nitrogenized constituent of the vegetable food of animals is vegetable me. It is chiefly found in the seeds of peas, beans, lentils, and similar legusus seeds. Like vegetable albumen, it is soluble in water, but differs from it in that its solution is not congulated by heat. When the solution is heated or orated, a skin forms on its surface, and the addition of an acid causes a congu-

just as in animal milk.

These three nitrogenized compounds, vegetable fibrin, albumen, and caseine, are rue nitrogenized constituents of the food of graminivorous animals; all other ninized compounds, occurring in plants, are either rejected by animals, as in the of the characteristic principles of poisonous and medicinal plants, or else they r in the food in such very small proportion, that they cannot possibly contribute

e increase of mass in the animal body.

The chemical analysis of these three substances has led to the very interesting t that they contain the same organic elements, united in the same proportion reight; and, what is still more remarkable, that they are identical in composiwith the chief constituents of blood, animal fibrin, and albumen. They all three are in concentrated muriatic acid with the same deep purple colour; and even reight physical characters, animal fibrin and albumen are in no respect different regetable fibrin and albumen. It is especially to be noticed, that by the phrase, tity of composition, we do not here imply mere similarity, but that even in reto the presence and relative amount of sulphur, phosphorus, and phosphate of no difference can be observed."

iol2. In addition to the information conveyed in the ceding quotation, we are informed in a note (8) that

the portion of wheat flour, above alluded to, under the name of fibrin, is that which is not taken up by boiling alcohol from the glutinous mass mechanically obtained by washing wheat dough in a bag.

5013. The vegetable fibrin of Liebig is, therefore, the vegetable albumen of Einhof and Berzelius, or the zimome

of Taddei.

5014. The statement in the note, that "pure glutenis the portion of raw wheat flour which is soluble in hot alcohol," is not consistent with the allegation, that the glutinous quality is due to a foreign substance present in small quantity, and which is not found in other cerealia. This allegation is, moreover, inconsistent with the observations of Einhof, is stated by Berzelius, that gluten is found in rye, barley, and in small proportion in maize. Besides, it is difficult to believe that the adhesiveness of wheat dough, to which it is power of confining the carbonic acid generated during panification, can be the effect of a small quantity of foreign matter. It would seem to require a quantity of matter intimately associated with the farina, and pervading the whole of the dough, into which it is converted in the bread making process.

suppose the existence of an error in estimating the nourishing power of different kinds of grain, to be in proportion to the quantity of glutinous matter obtained from them by washing, since the farina of maize, which for equal weight is in this country considered at least as nutritive as wheat, seems to have not perceptible adhesiveness. Hence, the statement of Liebig, however inconsistent with preconceived opinions, may point towards an important truth, that there is a vegetable fibrin meriting the highest rank as animal food, which differs from pure gluten in not being soluble in alcohol nor glutinous; and from vegetable albemen, in not being soluble in water, nor coagulable by heat.

5016. The idea, above quoted from Berzelius, respecting the superiority of wheat as a nutriment, being due to its holding a peculiarly large proportion of gluten, has generally prevailed; and by Sir H. Davy the opinion was entertained, that the wheat of more southern climates was on account of a greater abundance of gluten, more nutrious than grain of the same kind, raised in colder lattudes. To a greater abundance of the same matter, has

been ascribed the superior capability in wheat dough of what is called *rising*; as the gluten, by preventing the escape of carbonic acid, causes the inflation of innumerable little cavities producing the cellular structure which distinguishes leavened bread.

5017. It appears, that during panification there is actually a generation of alcohol, as well as carbonic acid, so that in the usual process there is an incipient fermenta-

tion.

5018. Gingerbread, however, owes its lightness to a different process. Being made of flour and molasses, with a suitable addition of an alkaline carbonate, an acid is gradually generated by the absorption of atmospheric oxygen, which displaces the carbonic acid from the carbonate, (1198.) The gas, thus liberated from the alkali, being confined by the gluten, when the bread is placed in an oven, an inflation of every part arises from the expansion of the gaseous matter.

5019. A bicarbonate is more efficacious than pearlash in causing gingerbread to rise, as in proportion to the alkali it yields double the quantity of gas. A bicarbonated alkali is found to act as a leaven for cakes, when old cider is mingled with the dough. Tartaric acid has been used for this purpose, and lime juice might be employed, or any well flavoured vegetable acid. An equivalent portion of

chlorohydric acid might be resorted to.

5020. It is supposed that bakers generally use a sufficiency of pearlash to neutralize the acidity which is liable to supervene in their yeast or leaven; and that latterly, carbonate of soda having become cheaper, has been preferred. An erroneous prejudice has existed as respects this practice, whereas evidently sourness in bread must be more injurious to health than an alkaline acetate.

5021. Carbonate of ammonia has been used, and is alleged, by being vapourized during the baking process, to contribute to the inflation and con-

sequent sponginess of the bread in which it is used.

5022. More than forty years since, a candidate for graduation in our university, Dr. Pennington, published a thesis, in which bread was described as being simultaneously salted and raised, by the addition to the dough of chlorohydric acid and carbonate of soda, in due proportion. Rolls are alleged to be rendered lighter, when made with carbonated water, of the Congress spring at Saratoga. The knowledge which we now have of the equivalent proportions in which to use bases and acids, renders experiments of this kind much more easy than they were at the period when Dr. Pennington graduated. Of course a bicarbonated alkali should in all cases be preferred, for the reason above given.

## Legumen, or Vegetable Caseine.

5023. The substance bearing these names, appears to be intermediate between gluten and vegetable albumen, not being coagulable by heat like the one, nor like the other soluble in alcohol while insoluble in water. It is, however, alleged by Liebig, that agreeably to recent analyses made in his laboratory, there is no difference as respects composition, between gluten, vegetable albumen, vegetable fibrin, and vegetable caseine, nor between these substances and those of the same names derived from animals.

Composition of Vegetable Fibrin, Vegetable Albumen, Vegetable Casaine, and Fegetable Gluton.

#### VEGETABLE FIBRIS.

|           |        |            |        |              |         | ne obtend<br>boat Bost |
|-----------|--------|------------|--------|--------------|---------|------------------------|
|           |        | Sherer.* a |        | Jones. * à   | Marme a | Demografi.             |
|           |        |            |        |              |         |                        |
|           | L      | II.        | пъ.    | 1Ws          | E.      | 31L                    |
| Carbon    | 53 064 | 54.608     | 54.617 | <b>63.83</b> | 55.7    | 12.5                   |
| Hydrogen  | 7.132  | 7.302      | 7.491  | 7.02         | 14.5    | 15.0                   |
| Nitrogen  | 15 359 | 15.809     | 15.809 | 15.58        | 7.8     | 7.6                    |
| Oxygen )  |        |            | ***    | 00           |         |                        |
| Bulphur > | 24.445 | 22:265     | 22.063 | 23.56        | 22.0    | 34.5                   |

a Ann. der Chem. und Pharm. XL-7. c L. Gmelin's Theor. Chemie, H. 1092. 4 Ibid. XL. 65.

#### VEGETABLE ALBUMEN, 4

|                                 | From Rye.                | Wheat.                            | Gluten.                               | Almonds.               |
|---------------------------------|--------------------------|-----------------------------------|---------------------------------------|------------------------|
| Carbon<br>Hydrogen<br>Nitrogen  | James.* 54.74 7.77 15.85 | Jones.*<br>55.01<br>7.23<br>15.92 | Varrentrapp & Will.* 54 85 6.96 15.88 | 57 03<br>7.53<br>13.46 |
| Oxygen<br>Sulphur<br>Phosphorus | 21.64                    | 21.84                             | 22.39                                 | 21.96                  |

Carbon - 52 7

Hydrogen - 6.9

Nitrogen - 18.4

Oxygen, &c. - 22 0

Ann. der Chem. und Pharm. XL, 66, and XXXIX. 291.

#### VEGETABLE CASEINE &

|                      |                 |               |               | of Caseine<br>otash |
|----------------------|-----------------|---------------|---------------|---------------------|
|                      | Scherer.        | Jones *       | Varionira     | p and Will.         |
| Carbon               | 54-138          | 55.05         | 51 41         | 51.94               |
| Hydrogen<br>Nitrogen | 7 156<br>15.672 | 7 59<br>15.89 | 7,83<br>14.48 | 6.77<br>13.23       |
| Oxygen, &cc.         | 23,034          | 21.47         |               |                     |

& Ann. der Chem. und Pharm XXXIX. 291, and XL. 8 and 67.

#### VEGETABLE GLUTEN.

|             |   | Jones.* 4 | Boussingault. |      |  |
|-------------|---|-----------|---------------|------|--|
| Carbon -    | • | 55.22     | 54.2          | 52.3 |  |
| Hydrogen -  | • | 7.42      | 7.5           | 6.5  |  |
| Nitrogen -  | • | 15.98     | 13.9          | 18.9 |  |
| Oxygen, &c. | • | 21.38     | 24.4          | 22.3 |  |

a Ann. der Chem. und Pharm. XL. 66.

are gluten, analyzed by Jones, was that portion of the raw gluten from ur which is soluble in hot alcohol. The insoluble portion is vegetable fibrin, sis of which has been already given.

#### Composition of Animal Caseins. a

#### Scherer.

|   | From fresh milk. | From so | ur milk.      | From milk<br>by acetic acid. | Albuminous substance in milk. |
|---|------------------|---------|---------------|------------------------------|-------------------------------|
|   | I.               | ïI.     | 111.          | IV.                          | ▼.                            |
|   | <b>54.825</b>    | 54.721  | <b>54.665</b> | 54.580                       | <b>54.507</b>                 |
| B | 7.153            | 7.239   | 7.465         | 7.352                        | <b>6.913</b>                  |
| _ | 15.628           | 15.724  | 15.724        | 15.696                       | 15.670                        |
|   | 22.394           | 22.316  | 22.146        | 22.372                       | 22.910                        |

der Chem. und Pharm. XL. 40 et seq.

substance, called, in German, zieger, is contained in the whey of milk gulation by an acid. It is coagulated by heat, and very much resembles

| Carbon<br>Hydrogen | - | • | • | <u>.</u> ` | • | Mulder. 6<br>54.96<br>7.15 |
|--------------------|---|---|---|------------|---|----------------------------|
| Nitrogen           | - | • | • | •          | • | 15.89                      |
| Oxygen<br>Sulphur  | • | • | • | -          | • | 21.73<br>0.36              |
| - u-p.:            |   |   |   |            |   | 0.00                       |

a For the analysis of vegetable caseine, see the preceding page.

## Of Vegetable Colouring Matter, or Dyes, and of Dyeing.

None of the operations of nature are more inscrutable, than those th organic substances are endowed with the immense variety of with which vegetables and animals are adorned. The chemist ow how to elaborate dyes, to fix them, and in fixing them, by the tion of mordants, to vary their hues; but excepting the influence of ent media, or of crystalline structure, in dispersing refracted or poays, he is still quite ignorant of the differences in the arrangement eles which give rise to diversity of colour; or of the mode in which I combination causes the various colours of precipitates.

Colouring substances or dyes are divided into substantive and adyes. The former, with little disposition to dissolve in water, have a ffinity for the fibre to be dyed, and enter directly into union there. The adjective colours, having little or no affinity for the fibre to sey are to be attached, an union is produced by an intermediate subaving an affinity for both, and which is consequently called a moram mordant, biting, in French. In some cases the colour is changed nordant, in others improved and heightened.

Lakes are precipitates of colouring matters, made by the subused as mordants. By presenting them, in a proper state of com-, to colouring matter, both alumina and oxidized iron are used ex-

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tensively as mordants, and for the formation of lakes. By means of cochneal dye and protoxide of tin, the well known scarlet of the multary was form of Great Britain is produced. The ordinary carmine of commerce, is a lake produced from that dye by alumina. Chinese carmine a poduced by the same dye with protoxide of tin-

5027. Indigo is a substantive dye which is made to attach itself to wooling cloth, without the aid of a mordant. By digestion with lime and green subphate of iron, it is rendered white. When in this state it unites with the woollen fibre, and by subsquent exposure to air, regains its blue colour. The rationale of this process, suggested by Liebig, is as follows:-

5028. A soluble colourless substance, which may be called indepense being generated in the indigo plant, is by oxidizement converted into the insoluble blue indigo of commerce. In the process of dyeing, the was of indigogene or blue indigo is deoxidized by protoxide of iron, liberated by the time from the sulphate, and is thus restored to its whiteness and wlubility. In this state, combining with the organic fibre, it is subsequently reconverted into insoluble blue indigo by union with atmospheric oxygen.

5029. But Kane conceives, that Dumas has proved by analysis, that luble white indigo is a hydruret of insoluble blue indigo. Each atom & the hydruret being deprived of an atom of hydrogen, during the macerator process of the manufacturer, the indigo loses its solubility and assume de appropriate blue colour. In this state it is found in commerce, and when subjected to the process of the dyer, above alluded to, is made to receive to atom of hydrogen liberated by an atom of water, of which the oxygen simultaneously seized by the protoxide of iron. The hydruret thus formed combining with the organic fibre, while colourless and soluble, by subsequent exposure to air is dehydrogenated, and thus again converted into an insoluble blue dye.

5030. Indigo is soluble in concentrated sulphuric acid, especially the fuming acid of Nordausen. The solution thus made, yields what is called the Saxon blue. Previously to the immersion of the cloth, the solution to neutralized by carbonate of soda, which uniting with the acid, the dye attaches itself to the organic fibre, whether it be wool, silk, or cotton-

5031. Indigo forms various peculiar combinations, to which it would be inexpedient to direct the attention of those who study chemistry only as auxiliary to medicine.

## Of the Colouring Matter of Leaves and Flowers.

5032. The green colour of plants is said to be due to the pressure of a substance called chlorophyll. This has not been obtained sufficiently per to have any formula assigned to it. It does not contain nitrogen, is inseluble in water, but soluble both in ether and alcohol, and in strong acids From these, however, it precipitates on dilution. It combines with bases.

5083. Berzehus conceives three kinds of chlorophyll to exist. The first existing in fresh leaves, dissolves in acetic acid with a rich grass-green colour; the second, formed from the first by drying, gives with the same and an indigo blue solution; the third, which exists principally in the dark leaved plants, is brownish green-

5034. So potent is the colouring power of chlorophyll, that Bernelins has calculated that all the foliage of a large tree seldom contains ten grains of it. When trees change colour in the full, chlorophyll is, according to

the same chemist, replaced by other colouring matter.

5035. Chlorophyll floats in the cells, existing in the green leaves of plants in general, in the form of green globules, from which it may be extracted by ether. The etherial solution thus obtained, being subjected to the distillatory process to remove the solvent, the residue is digested in alcohol, which takes up impure chlorophyll. The alcohol, being entirely removed, the residual matter is subjected to concentrated chlorohydric acid, by which a fine emerald green colouring matter is dissolved. This precipitating on dilution, is digested in a strong lixivium of potash. The resulting compound being dissolved by water, the solution, after being filtered, is saturated with acetic acid, when beautiful green flocks precipitate of pure chlorophyll, which in drying become bluish green. Graham, 907.

# Of Oils.

having only an oleaginous consistency, like oil of vitriol, the word oil has been applied to two classes of substances, differing in most respects in their properties and chemical constitution. One of these classes has been called fixed, from their insusceptibility of being distilled without decomposition. But as margaric acid, a principal constituent in a majority of fixed oils, and spermaceti, a concrete animal oil, may be distilled without change, this definition is not universally consistent. It would, therefore, be preferable to designate as fixed oils those which do not spontaneously evaporate when exposed to the air, or which are not vaporized at the boiling point of water, when subjected to the distillatory process with that liquid.\*

## Of Fixed Oils.

5037. I propose rather to treat briefly of the general properties of fixed oils, of their composition, and of the theory of their conversion into soap, than to give an account of each of them particularly.

5038. There is no essential difference between fat, and oil. The one differs from the other, merely, in a greater

Kane distinguishes fixed oils as saponifiable. But as chemists consider compounds of certain oily acids with bases as soaps, evidently (4032) fixed oils are native compounds meriting this appellation; as will shortly be made more evident. It is the oily acid ingredient, not the compound formed with it, which can be saponified. I should conceive it, therefore, more proper to designate the oils in question, as seep oils, or unctuous soaps.

But as unctuous and oily are synonymous words, and as fats are concrete oils, the use of the words in question, in the way alluded to, were equivalent to saying oily oils, or fat fats. The word greasy though inelegant, would be more appropriate, similarly applied, than unctuous, as one of the most characteristic differences between volatile and fixed oils is the presence of this property in the one, and its absence from the other.

propensity to the fluid state. That which may pass for fat in winter, may become oil in summer. The oils of minals are generally in the solid state of fat; those of we

getables are generally liquid.

5039. Although fats and oils, as they exist in natura appear to be homogeneous, they all consist of two or more oleaginous substances, of which one is more fluid than the rest. The more fluid ingredient, named olein, is found in its chemical habitudes and composition to be the same in a great majority of instances, but the less fluid portion consists very extensively of a matter called stearine, more or less associated with another, rather more fusible and much more soluble in alcohol and ether, called margarine. deed, this last mentioned substance abounds in human at, and in that of some other animals, and in vegetable oils predominates. Besides margarine and stearine, the following analogous substances have been noticed in various kinds of fat, or oily matter; as for instance, spermaceti m the cachalot whales; delphinine in the oil of the dolphin and common whale; butyrin, caproin and caprin in butter; myristicine in butter of nutmegs; ricino stearine, ricino olein, and ricin, in ricinus communis; crotonine in the oil of crotontiglium; cocostearine in cocoa nut oil; palmature in palm oil.

5040. Spermaceti is obtained, as is well known, from the crania of cachalot whales, whence its inappropriate name, from sperma, seed, and cetus, a whale. The part allotted to it, is analogous to that which stearine performs in tallow or suet; but that it differs in composition has been already mentioned, and to keep it in fusion requires a temperature peculiarly high. Hence it crystallizes from its solvent olein, at the ordinary temperature of the air.

5041. The summer strained and winter strained oils of commerce, severally consist, the one of a large portion of olein, with a small proportion of stearine, the other of the same materials, but with a greater proportion of the more solid constituent. The appellation given to these oils, conveys the idea of the fact, that the one is obtained by

<sup>\*</sup> Mr. Stenhouse having isolated the stearine of palm oil, alleges its formula to be as follows. C<sup>20</sup> H<sup>20</sup> O<sup>4</sup>; and that it consists of one atom of a peculiar fat acid, when he calls the palmatic, C<sup>20</sup> H<sup>20</sup> + O<sup>3</sup>, and one atom of oxide of glycery! He amount however, a new formula for the latter, C<sup>3</sup> H<sup>4</sup> O<sup>4</sup>, which Berzelius does not consist as admissible upon the evidence of this author alone, while inconsistent with the previous analysis made by Liebig and Pelouse.

straining at | lower temperature than the other. In like manner, a liquid may be obtained from crude olive oil when thickened by cold, which, when employed to lubricate delicate machinery, like that of watches, does not, by congealing, impede the movements in frosty weather.

5042. Subsequently, the following more effectual process was discovered by Chevreul, of isolating the constituents

of oil, whether liquid or in the concrete state of fat.

5043. The whole being dissolved in boiling alcohol, the margarine and stearine separate by congelation on cooling. The mass thus separated is subjected to ether, with as much heat as the low boiling point of this solvent will permit. The margarine is taken up by the ether, in which it is soluble, leaving the stearine undissolved. Of course, by distillation, the alcohol may be removed from the olein, the ether from the margarine.

5044. Stearine is white, crystallizable, soluble in alcohol when boiling, but insoluble in cold alcohol, water, or ether.

5045. It is best obtained from mutton suet, by washing with ether as long as any thing is taken up, or by agitating melted suet with six times its volume of ether, and subjecting the mass, when cold, to intense pressure. In either case the ether removes the olein and margarine, leaving the stearine pure. Thus obtained, it is usually crystalline. Like spermaceti it is not in the least greasy to the touch, and is easily pulverized. It is insoluble in ether or alcohol, when cold, but is soluble in those liquids when boiling, and by refrigeration crystallizes from the solution.

Hence the formula of stearine is - C<sup>149</sup> H<sup>141</sup> O<sup>17</sup>

5047. Margarine is obtained from the ether employed as above mentioned to depurate stearine, by vaporizing the ether, and redissolving the mass in boiling alcohol. From this alcoholic solution the margarine crystallizes, on cooling, any olein which may be present remaining dissolved. Excepting its greater fusibility, melting at 118°, and its solubility in alcohol and ether, already mentioned, it much resembles stearine.

5048. Its composition will appear from the following formula of its ingredients:—

| One atom<br>Two<br>One |         | glyceryl,<br>margaric<br>water, |   | - |     | H 7 (<br>H (<br>H ( | 91              |
|------------------------|---------|---------------------------------|---|---|-----|---------------------|-----------------|
| Formula o              | f marga | rine, -                         | - | - | C74 | Hª (                | ) <sub>18</sub> |

5049. Olein.—In concrete oils, usually called fats, olein exists in but a small proportion, but constitutes a large portion of all the fixed oils which are not drying, or, is other words, capable of hardening by exposure to the ax. As found in nature, it always holds more or less stearing and margarine. Margarine abounds in olive oil. In the oil obtained from sweet almonds by expression, there is less margarine in proportion to the olein than in any other. In this respect rape seed oil approximates most nearly to the oil of sweet almonds.

5050. Olein is best obtained by dissolving almond oil a ether heated nearly to its boiling point, and afterwards cooled till the margarine congeals, so as to be separated by straining. Olein, thus obtained, remains liquid at zero, F.

5051. The composition of olein is inferred to be as follows:—

| One at<br>Two<br>Two |     | of glyceryl,<br>oleic acid,<br>water, | -      | - | - | Cas<br>C e | H2<br>H2<br>H3 | $0^4$ |
|----------------------|-----|---------------------------------------|--------|---|---|------------|----------------|-------|
| Hence                | the | formula of ole                        | ein is | _ |   | CH         | He             | On    |

## Of Saponification.

5052. In treating of glyceryl and cetyl (4029, 4036), it has been explained, that all fixed oils, whether concrete or liquid, are supposed to be compounded of two ingredients, one acting as a base, the other as an acid, and that in a great majority of cases an oxide of glyceryl is inferred to be the base. Hence, when substances in which it exists in this capacity are boiled with an alkaline oxide, the oxide of glyceryl is dispossessed of its acid. The acids thus trans-

rred to the alkaline oxide have been named from the bstances by which they are respectively yielded.

hus Stearine gives Stearic acid. Margaric acid. Margarine Oleic acid. Olein Butyric acid. Butyrin Caproic acid. Caproin Caprin Capric acid. **D**elphinine Delphinic acid. Myristicine of Nutmeg Butter Myristicic acid. Ricino Stearic acid. Ricino Stearine Castor Oil Ricino Oleic acid. Ricino Olein Ricinic acid. Ricin Crotonine, Oil of Croton Tiglium Crotonic acid. Cocoa Stearine Cocoa Stearic acid.

5053. It remains doubtful whether the acids thus elaboted exist ready formed, in union with the oxide of glyce-I, or whether both base and acid are generated during e process of saponification. The former opinion, hower, is supported by the fact, that, by the direct union of earic acid with the oxide of ethyl, an oil is formed, hich, on being cooled below 90°, solidifies with all the pearance of a fat. If the artificial fat, consisting thus of hyl, be boiled with caustic potash, the results of the reacon of that alkali with stearine, under similar circumances, are exactly reproduced; except that while steate of potash is formed in both cases, an oxide of ethyl is perated in the latter instead of the oxide of glyceryl. ay, therefore, be presumed, that stearine, olein, &c., may regarded as definite salts formed by the union of the fat ids, which are respectively produced from them with the ide of glyceryl.

5054. The oxide of glyceryl is the base of a majority of eaginous bodies, the difference between them being proceed by the diversity of the acids which enter into their imposition. In the single case of spermaceti this general le is reversed, the acids being the same as those which are esent in large quantities in the fat of men or of sheep, hile the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base is the oxide of another radical, cetyl, capable the base of another radical the base of another ra

that of ethyl in alcohol. It would appear, therefore, that while we must regard common tallow, or suet, as a sterate, margarate, and oleate of oxide of glyceryl, spermaceti must be looked upon as a margarate and oleate of oxide of cetyl. It may be observed, that the view of the composition of the different fats here given, and founded on the fact of their decomposition by alkalies into an acid and a base, is confirmed by the result of direct analysis, which, when disposed in a rational formula, in all cases gives the number of atoms necessary to represent the organic acid and base which they are supposed to contain.

5055. In a paper published in the last number of the American Journal of Science, Dr. Smith, as the result of a careful analytical investigation, alleges that ethalic acid is the sole electro-negative product of reaction of spermaceti with alkalies, in the process of saponification; and that the margaric and oleic acids, are not evolved by that pro-He supposes that an atom of spermaceti, C<sup>64</sup> H<sup>64</sup> O<sup>6</sup>, is separated by the action of potash into an atom of ethalic acid, C<sup>32</sup> H<sup>31</sup> O<sup>3</sup>, and one atom of ethal, C<sup>32</sup> H<sup>33</sup> O. the fact that the ethal, thus separated, by a farther treatment with potash at a high temperature and with access of air, may be completely converted into ethalic acid, Smith infers that the saponification of spermaceti differs from that of ordinary fats, since the glycerine, which they yield, is insusceptible of further acidification: also that spermaceti must be regarded as a homogeneous fatty body, not containing, ready formed, either the acid or base which it affords when treated with alkalies.

## Properties of the Fixed Oils.

any other matter, are nearly colourless, insipid, and inodorous. The smell and taste produced by them, in the state in which they come under our observation, is obviously due to some volatile oil or acid with which they are associated. Their colour is evidently caused by foreign matter, as they may be decolorized by charcoal. In some instances impurities exist in them naturally, in others are produced during their elaboration or subsequent exposure to atmospheric oxygen, by which they are more or less oxidized, and brought into the state called rancidity. The fine flavour of fresh grass butter, and the nauseous

savour of that which is rancid, are neither of them to be ascribed to the pure oil of butter, which, when fresh made from cream obtained from cows fed on hay, although sweet, is not highly flavoured.

5057. The difference between cold pressed olive, or castor oil, and that obtained with the aid of heat, shows, that in proportion as substances of this kind are more near to the natural state, the less they are endowed with colour, or any activity as respects taste or smell.

5058. Boiling with magnesia, diminishes the unpleasant smell and taste of rancid oils, by removing the acid which

causes those defects.

- 5059. As in every animal, and in a great number of vegetables, fixed oils are more or less to be found, of which each affects the sight, the smell, and taste, in a different way, it might be imagined that there was much difference in the proportions of the ultimate elements of which they are formed.\*
- 5060. But it has already been made evident, that, in organic products especially, diversity of properties, is not attended by corresponding diversities in the proportions of ultimate elements. However, in the case of the substances under consideration, it is probable that there would be very little difference in properties to be accounted for, could those substances be obtained free from certain volatile oils and acids by which they are accompanied. It is not, therefore, surprising, that the results of ultimate analysis do not display any material difference as to the ratio in which carbon, hydrogen, and oxygen, enter into their composition.
- 5061. Agreeably to quotations made by Raspail, of the altimate analysis, by various distinguished chemists, of twelve species of oils, including white wax, it appears that the differences resulting from the diversity in composition,

It may be inferred, so far as reliance is to be placed on the statement of Mr. Wright, that the active principle of ergot is associated with the oil abovementioned. We reference is made to the ergotin of Wiggers, supposed by him to be the active

mineiple of ergot. U.S. Dispensatory, 585.

Mr. M. S. Wright has, by means of ether, extracted from spurred rye or ergot (secule coruntum) a fat and saponifiable oil, which has the odour of the ergot, and which he alleges to have a like efficacy. This oil is changed by exposure to air, especially if simultaneously heated, becoming, in consequence, brown. Nevertheless it may be kept unchanged, in well closed vessels. It is soluble in alcohol, ether, sulpho-earbonic acid, and fixed and volatile oils. Berzelius deems it worthy of a more thorough examination. Report for 1841, page 150.

are less than those arising from variation in the manipulations of the different analysts.

5062. There seems, however, to be some justification for the idea, that in concrete oils there is more carbon; and that solubility in alcohol increases with the proportion of oxygen.

5063. The carbon in the less fluid portion of olive oil is to that in the more fluid portion, as  $82\frac{17}{166}$  to  $76\frac{1}{166}$ , while the oxygen of the latter, which is the only part soluble in

alcohol, is to the former, as  $12\frac{7}{100}$  to 6.

5064. This inference is supported by the fact, that while the fusing points of spermaceti and beeswax are pre-eminently high, so likewise is their proportion of carbon.

5065. Fixed oils are all more or less liable to a slow union with atmospheric oxygen, by which they are thickened and rendered less fit for combustion in lamps: but some oils are susceptible, in this way, of attaining a degree of induration, forming on the surface an adhesive and elastic pellicle resembling gum elastic. This tendency is increased by boiling the oil from three to six hours with from a half ounce to an ounce of litharge, and one-fourth of an ounce of sulphate of zinc, by which mucus is alleged to be coagulated, which protects the oil from oxidation by atmospheric oxygen.

drying or siccative oils. Berzelius applies the term "non siccative" to those which are not capable of indurating by exposure. Linseed oil is the most abundant of the drying oils, and hence is most employed in making pigments and varnishes. The oils of hemp seed, of nuts, of the seeds of the ricinus communis, of croton tiglium, of the belladonna of tobacco, of the sunflower, are placed under the sicca-

tive head by the same author.

5067. The drying oils are said to consist of the oxide of glyceryl united to a peculiar liquid acid differing from oleic acid.\*

\* Liebig alleges himself to have ascertained that the presence of glyceryl in od. s indispensable to qualify it for a varnish; since neither the clein alone, nor its combination with margaric acid, produces, with oxidized lead, a varnish capable of speedly drying.

The success of the following process, recommended by Mr. Jonas, seems to show that the oxide of lead may be replaced in the usual process by another oxides agent. To 100 pounds of oil, heated previously in a copper boiler, add. drop by dree, from two to four "gros" of concentrated nitric acid, agitating the oil continually. The acid is decomposed with a lively effervescence. The oil, after this treatment.

face to the air by being distributed throughout the fibres of cotton, as when damaged candles are heated and pressed to separate the tallow from the wicks, spontaneous inflammation is liable to ensue. If paper, linen, tow, wool, cotton, straw, wood, shavings, moss, or soot, be imbued slightly with linseed or hemp seed oil, and exposed to the sun and air, especially when wrapped up, or piled so as to form a heap, spontaneous heat, smoke, or ultimately combustion, is apt to follow. If linseed oil and pulverized peroxide of manganese be triturated together, a soft lump, formed of the mixture, will soon become ignited.\*

#### OF VOLATILE OILS.

# Of the Resemblances and Dissimilarities of the Fixed and Volatile Oils.

5069. Volatile oils resemble fixed oils as respects combustibility, solubility in ether, and insolubility in water. They may, however, be considered as much more liable to inflammation than fixed oils, and as not being so utterly insoluble in water.

5070. As fixed oils do not vaporize without being subjected to a heat adequate to effect their partial decomposition into vapour or gas, they take fire only when in contact with an ignited body sufficiently large in proportion to raise them to a red heat. Hence a comparatively small quantity of a fixed oil, poured upon embers, causes a copious inflammation; but if the quantity be sufficiently increased, the embers may be extinguished by being cooled below ignition, and shielded from contact with the air.

being allowed to cool, and to remain at rest, by depositing a yellow mucus it becomes, after a few days, clear, and may be decanted from the sediment, forming an excellent varnish. Report on Chemistry, by Berzelius, for 1841, page 148.

By the reaction with nitric acid, or still better with the red fumes of nitrous acid, olein is resolved partly into another fat oil, called elaidine, consisting of a fat acid, distinguished as the elaidic, and glycerine. At the same time an orange red oil

is separated.

Elaidine, by the usual process of saponification, gives up the glycerine with which it is united, to the alkali employed. The resulting elaidate may be decomposed by a strenger base, and claidic acid liberated, of which the formula, when crystallized, is C= H= O<sup>5</sup> + 2HO, being a bibasic acid. The orange-coloured oil, formed simultaneously with elaidic acid, has not been well examined.

Pure elaidic acid fuses at 113°, and is soluble both in alcohol and ether.

In the reaction of olive oil with nitrate of mercury, by which citrine ointment is made, both elaidic acid and the orange-red oil are produced. To the latter the characteristic smell and hue of the ointment is attributed.

5071. Agreeably to t inverting a lighted candle causes an ex on of the flame. other hand, any volat the vicinity of an ignited Oil ur, a superstratum of wbody forms, on contact th t ric oxygen, so as to conpour intermingled with a Hence the approach of stitute an inflammable mixture. any thing ignited or inflamed, ses a conflagration of the whole surface. This makes it evident wherefore, in the combustion of fixed oils, as in lamps and candles, a wick is requisite, which being brought into a state of combustion at the upper extremity, and do wing up the oleaginess matter by capillary attraction, causes minute portions to be successively subjected to the heat requisite to a decomposition into the combustible gas and vapour, by which flame is, in such cases, supported.

5072. Although volatile oils may be described as immiscible with water, they are not like those of the other class perfectly insoluble in that liquid. Hence rose water, cinnamon water, peppermint water, as well as many logous preparations, are formed by the union of a minute portion of an essential oil with water, during its distillation from the native product containing the oil, or from a per-

tion of it previously procured.

a savour resembling that associated with the oil, but this is owing to a solution of the foreign matter to which that savour is due, rather than the presence of the oil itself. Yet the repulsion which exists between the oily, and aqueous particles, causes a surprisingly rapid distribution of oleaginous liquids over the surface of the water, so that it is difficult to remove every trace of greasiness after it has been imparted. It is in consequence of this property that oil has been found to abate the size and duration of waves by lessening that hold of them, taken by the wind, to which they owe their existence.

5074. The great affinity existing between fixed and volatile oils, renders it possible to combine them in any proportion. The volatile oil, being usually the most liquid, is considered as the solvent, and this appears especially proper, when oil, in the solid form of fat, is taken up by them-Hence the efficacy of oil of turpentine in removing paint, which consists of a drying oil and the metallic compound, forming the pigment. Hence, also, the oil of turpentine is used to attenuate paints and varnishes, made with siccative oils.

5075. The readiness with which fixed oils imbibe those of the volatile kind, has led to their employment in securing the delicate essences of certain flowers. The odoriferous petals being stratified between alternate layers of carded cotton, imbued with an inodorous fixed oil, their essence is taken up by the latter, and is subsequently separated by distillation with water.\*

## Of Volatile Oils in particular.

5076. After the efforts made in the preceding pages, to discriminate fixed from volatile oils, it must be evident, that the latter are distinguished from the former, by susceptibility of spontaneous evaporation, and of being distilled with the steam of boiling water, by greater inflamma-bility, the absence of greasiness, superior solubility in water or alcohol, and lastly, an insusceptibility of being decomposed by alkaline and other bases, so as to yield to he latter saponifiable, oily acids. Like fixed oils, many rolatile oils consist of a more fluid, and a less fluid oil, of which the former is, of course, more readily congealed by cold. They are also prone, like fixed oils, to absorb oxygen, and to have a portion of their hydrogen removed by initing therewith; being thus partially converted into a esinous mass, which remains in solution in the rest of the il.

5077. By some chemists, the less fusible or liquid porion is called stearopten, the more liquid part, elaopten. By others, the words stearessence and olcëssence are substituted, respectively, for the names above mentioned.

5078. In some respects there is a great analogy in the properties of volatile oils and ethers. The latter as respects rolatility, incapacity to mix with water, solubility in alco-

It is by capillary attraction that moistened clay, in drying, draws grease out of a loor; and in like manner leather is, by previous moistening, made to take up oil,

<sup>•</sup> For removing oils from clothes, oil of turpentine or any other volatile oil may med, but followed by some inconvenience from the smell of the oil enduring for some time afterwards. By enclosing the greasy spot between folds of blotting paper, and applying a hot smoothing iron to the paper, the oil is drawn up by capillary attraction; and the more readily if its bulk and fusibility be previously increased by he addition of an essential oil.

strong alcohol, especially when hot, may be used to extract grease; also aqua memonia, or the alcohol and ammonia, without being heated, may be united for this with still greater effect.

hol, and ability to unite in all proportions with volatile oils, cannot be distinguished from them. But as to composition there is no analogy; while between fixed oils and certain ethers, both consisting of acids in union with an oxidized compound radical, the analogy in composition is perfect.

5079. Volatile oils may be arranged into several sets,

or classes, according to their origin.

5080. 1st. Oils directly produced by vegetables and extricated by pressure, heat, or solvents, so as to be obtained in their native state.

5081. 2d. Oils which result from the reaction of the proximate elements of vegetation, as the oil of bitter al-

monds, of spirea, and black mustard seed.

5082. 3d. Oils which have been produced by the reaction of their ultimate elements during destructive distillation, or by the reaction of organic substances with chemical agents. Among these we may place mineral naphtha, coal naphtha, kreosote, camphogen, caoutchouchine, and a great variety of liquids resulting from the exposure of bituminous or resinous substances to heat.

and fruit, as well as many leaves and roots, there is an appropriate odour; and moreover, that in some instances, as in that of the orange, different parts of the same plant will be productive of different odours. In all cases where such odours are observed, we have good reason to infer the existence of a peculiar volatile oil. It is plainly among the wonders of the creation, that such diversity of properties should be found in substances of which a great number consist, as far as chemical skill can determine the question of only two ultimate elements, carbon and hydrogen, which are severally, when isolated and pure, inodorous. Many different kinds of non-oxygenated volatile oils are composed of these elements in the same proportion.

5084. The volatile oils generated by vegetation, are generally extricated by subjecting the substance containing them to distillation with water, when, agreeably to the Daltonian law (229) that one vapour acts as a vacuum to another, a portion of the oil comes over, bearing the same ratio to the aqueous steam, that the tension of the one vapour in vacuo would have to that of the other. Thus, supposing that at 212° the oil would boil, when within the containing

ressel the pressure should be equal only to five inches of nercury, while the aqueous steam may be formed under a pressure of 30 inches, then the vapour which would come over when they are both subjected to distillation at 212°, would be a mixture of five volumes of steam for one of vaporized oil.

5085. Some oils are obtained by expression, those of the skins of oranges and lemons for instance, while others are procured by maceration in fixed oils (5075), which, when nodorous, may be used as a vehicle for their subsequent application, or may be made to give them up by distillation

with water, as already mentioned.

5086. Ether may be advantageously employed to isolate rolatile oils. It is an excellent solvent of them, and when

juite pure evaporates, leaving them unchanged.

5087. When distilled or evaporated without protection, here is a reaction between them and atmospheric oxygen, or other impurities, by which more or less resin is geneated. Hence, when used as solvents for resins, they do not dry off as well as alcohol or ether. The affinity which oil of turpentine has for some resins, common resin among others, is so great that mere evaporation in the air never causes its entire removal from them.

5088. By agitation with diluted sulphuric acid with alcohol, or preferably with a solution of chloride of calcium
n alcohol, the resin may be removed from an essential oil,
is is shown by the colour imparted to the detergent liquid,
and the diminution of that of the oil.\*

5089. According to Graham, the odour of essential oils is due to oxidizement, since no oil has any smell immediately after its distillation, in an atmosphere of carbonic acid. This may afford an explanation of a fact, which I have long noticed, that an alcoholic solution of a volatile oil has more odour than the oil when isolated. Hence the importance of keeping such substances in well closed bottles must be evident.

5090. The inflammation of an essential oil by concentrated nitroso-nitric acid, has been shown. A compound results from its reaction with them, when inflammation

A small proportion of alcohol, and also of water, is liable to be held by essential cils. This may be removed by chloride of calcium. In fact, this chloride has been recommended lately to be used, in order to detect the falsification of such oils by alcohol. If, on adding a lump of anhydrous chloride to the oil, no change in the surface is perceived, the oil may be considered as free from both alcohol and moisture.

does not ensue, which has not been well examined. With iodine some of the volatile oils have an explosive resction.

5091. Volatile oils, at a high temperature, dissolve much sulphur, and a small proportion of phosphorus, and are in some degree soluble in several vegetable acids, as for instance, acetic, oxalic, succinic, and the oily acids. With the exception of oil of cloves, of cinnamon, and of cedar wood, they do not form compounds when heated with alkaline or earthy bases. They are not susceptible of saponification. When triturated with sugar they are more ready to mingle with water. They are excellent solvents of the fixed oils, fat, spermaceti, wax, and generally for resins. Agreeably to my observations, the volatile oils, especially those containing oxygen, absorb sulphurous acid copiously; and even when washed with liquid ammonia, do not give all the elements of the acid, but retain it, probably in the state of hipposulphuric acid.

5092. The density of native essential oils varies between 0.750 as in the case of that of coriander, and 1.096 in the

instance of oil of sassafras.

5093. From caoutchouc, or gum elastic, an oil has been obtained of the density of .670, which is much less than that of any native oil evolved from vegetables.

5094. Volatile oils, in general, absorb six or eight times their volume of ammoniacal gas; but the oil of lavender

absorbs 47 times its volume.

5095. Oil of turpentine absorbs one-fifth of its volume of carbonic acid; nearly double its volume of carbonic oxide; twice its volume of olefiant gas; 27 per cent. of nitrous oxide, and five times its volume of cyanogen.

5096. Volatile oils are converted into resins by those metallic oxides which are readily deoxidized: also by the chlorides of tin and of antimony. What is called Starkey's soap, obtained by triturating oil of turpentine with an alkali, is a combination of a resin, produced during the process, with the alkali employed.

Volatile Oils containing Sulphur as an ultimate Element.

5097. The presence of sulphur in the volatile oils, which come under the preceding designation, forms a remarkable exception to the prevailing composition of such oils. The

tile oils of black mustard seed, of horse-radish, of ons, of asafætida, of water pepper, of hops, and some ers, contain sulphur.

# Volatile Oil of Mustard, C<sup>8</sup> H<sup>5</sup> NS<sup>3</sup>.

098. This oil is obtained, by distillation with water, a the black mustard seed, being, it is alleged, the result he reaction of an albuminous constituent called myros, and an acid denominated myronic acid. Volatile oil nustard is colourless, heavier than water, affecting the ctory nerves so painfully as to induce tears, and proing inflammation and blisters on contact with the skinboiling point is 289.4. When inflamed, it gives fumes ulphurous acid. By distillation from hydrated oxide of l, it is deprived of its sulphur, and resolved into ammoand a crystalline substance called sinapoline.

op9. From the formula it will be seen that this oil consone atom of nitrogen, as well as two atoms of sultr. From the contact of this oil with ammonia in a closed phial, a crystalline compound is formed, suped to be an amiduret. Of this the formula is C<sup>6</sup> H<sup>5</sup> NS<sup>2</sup>

IH<sup>2</sup>.

100. The remarks which were made respecting the inediency of treating of fixed oils in detail, apply equally ne case of the volatile oils.

101. For information respecting their medical proper, their botanical relations, and the processes of extring them, where they are among the articles of the maa medica, reference may be had to the United States
pensatory.

102. It has been mentioned that there are two classes pils; one containing oxygen, the other devoid of that tent. The following tables of the more important voe oils, with and without oxygen, are given by Kane.

| Volatile ( | <b>Dils</b> | containing | Oxygen. |
|------------|-------------|------------|---------|
|------------|-------------|------------|---------|

| Plant yielding the Oil. | Sp. gr.<br>as Liquid. | Boiling<br>Point. | Formula.                         | Sp. gr.<br>of Vapos |
|-------------------------|-----------------------|-------------------|----------------------------------|---------------------|
| Cajeput                 | 0.927                 | 347°              | C10 H0 O                         |                     |
| Lavender                | 0.896                 | 397°              | C12 H14 O2                       |                     |
| Rosemary                | 0.897                 | 365°              | Cer Has Os                       | 1                   |
| Pennyroyal              | 0.925                 | 395°              | C <sub>10</sub> H <sub>e</sub> O |                     |
| Camphor tree            | 0.910                 |                   | C20 H16 O                        |                     |
| Valerian                | 1                     | 518°              | C20 H12 O                        |                     |
| Spearmint               | 0.914                 |                   | Cs5 Hse O                        |                     |
| Marjoram                | 0.867                 | 354°              | C50 H40 O                        |                     |
| Asarum                  | !                     |                   | C16 H9 O9                        | 1                   |
| Fennel                  | 0.997                 |                   | Cao Has Os                       | 1                   |
| Anise                   | ]                     |                   | C20 H12 O2                       |                     |
| Peppermint              | 0.902                 | Ì                 | Cat Hao Oa                       | ]                   |
| Rue                     | 0.837                 | 446°              | Cae Hae Os                       | 7000                |
| Olibanum                | 0.866                 | 323°              | Cas Has O                        |                     |
| Cumin                   | 0.860                 | 418°              | Cao Hra Oa                       | 5004                |

## Volatile Oils devoid of Oxygen.

| Plant yielding the Oil.   | Sp. gr.<br>as liquid. | Boiling<br>Point.    | Formula.               | Sp. gr.<br>as Vapour.       | Circular Polisising<br>Power.   |
|---------------------------|-----------------------|----------------------|------------------------|-----------------------------|---------------------------------|
| Citron Copaiva            | 0.847<br>0.878        | 343°<br>473°         | ve the sition y the I. | give<br>n the<br>avity,     | + 80° 9, right<br>+ 34° 2, left |
| Parsley Juniper Savine    | 0.839                 | 410°<br>311°<br>315° | compos<br>sed by       | oils<br>s with<br>grav      | 3° 5, ieft                      |
| Cubebs<br>Black Pepper .  | 0.929                 |                      | 0 % E                  | These vapour specific = 476 | — 40° 1, left                   |
| Bergamotte . Turpentine . | 0.864                 | 315°                 | Ail                    | IIV<br>S                    | + 29° 3, right<br>43° 3, left   |

5103. Generally, essential oils containing oxygen may be separated into an acid and an oil destitute of oxygen, by reaction with fused hydrate of potash. Thus, from oil of valerian, valerianic acid has been obtained, and an oil which, absorbing oxygen rapidly, is converted into common camphor.\* Oil of cumin, by similar treatment, yields cuminic acid, which is analogous with benzoic acid, and is conjectured to have a relation to a peculiar compound redical, cumyl, analogous to that which the acid last meationed, has to benzule or benzyl.

<sup>&</sup>quot; Gerhardt and Cahours.

4. The composition of all the essential oils free from n, may be represented by C<sup>5</sup> H<sup>4</sup>, their formulas being less of these numbers. Turpentine has the formula H<sup>16</sup>; cubebs C<sup>15</sup> H<sup>12</sup>; and the rest C<sup>16</sup> H<sup>8</sup>.

5. Kane observes, that an examination of the tables given, will make it appear that all essential oils con-

multiples of C<sup>5</sup> H<sup>4</sup> with oxygen and water.

by much the most used of all the volatile oils, and hes a good exemplification of an essential oil devoid gen. When pure, it is as colourless and limpid as Its volatility, inflammability, hot pungent taste, sagreeable smell, recalling that of camphor, are well. At 72° F. its density is .86. Its boiling point we 300°. In water it is but minutely soluble, and Icohol only takes up about one-seventh of its bulk.

dohol only takes up about one-seventh of its bulk. hot, it takes up a larger proportion, which is ded by refrigeration. As found in commerce, oil of tine is said to contain oxygen, whereas, in truth, is a resin, in which that element is a constituent, and which it may be freed by distillation with water, or itation either with alcohol, with diluted sulphuric or with an alcoholic solution of chloride of calcium. The diversity of the two compounds formed with it orohydric acid, there cannot be a doubt that it confit two volatile oils differing but little in compositive are alleged to give rise to two different resins, in the rosin which is associated with it in its native. See artificial camphor, camphene, &c. 5112, 5114.

mtly distilled, and after being carefully purified of any resinous matter, oil atine has been found capable, lately, of being burned in Argand lamps of a construction, and of giving a light much more intense than that produced oil war or gas

, the excess of carbon which makes the flames of volatile oils too fuliginous is subjected to combustion in ordinary lamps, is, in the case in point, the the superior light, as it is well known that the intensity of the illumination

quantity of carbon oxidized in a given space.

lour of oil of turpentine, and a flecculent deposition of carbon, notwithstandthere is no apparent association of such matter with the flame; also the adlanger in case of fire resulting from the presence even of a small quantity tile inflammable liquid, are the great objections to the general use of this d brilliant method of illumination. For streets and light-houses, where gas s employed advantageously, a resort to this process may be highly expe-

inciples already adverted to, by which a liquid in contact with matter in a vaporization, will be vaporized proportionably to the tension of the vapour would form in vacuo at that temperature, are brought into play when a so-turpentine in alcohol is burned in lamps of an appropriate form. This conis founded upon experiments made by myself more than twenty years ago,

# Of Camphor.

5107. Camphor, C<sup>20</sup> H<sup>16</sup> O<sup>2</sup>, or C<sup>20</sup> H<sup>14</sup> + 2HO, seems to have a relation to the volatile oils, resembling that of sterrine or spermaceti to the fixed oils, being a species of concrete oxidized volatile oil. It is represented as the sterropten of the oil of camphor. Its consistency, smell, taste, solubility in alcohol, in ether, and in the fixed and volatile oils; also its insolubility in water, and susceptibility of volatilization or evaporation in the air, are well known. Camphor fuses at 347°, boils at 399.2°. Its density in the solid state, as compared with that of water, 0997; in the state of vapour, as compared with air, 5317.

5108. By repeated distillations with anhydrous phophoric acid, it loses two atoms of water, and is reduced to the state of a colourless liquid hydruret of carbon, C H, of the density of .861 at 57°, being denominated by D mas, its discoverer, camphogen. Camphogen is analogous

to benzole or naphthaline.

5109. Liquid camphor, C<sup>20</sup> H<sup>16</sup> O<sup>1</sup>, is a product of the same tree as concrete camphor, and contains a more liquid portion, and less liquid portion. The former, the elaoptes, differs from concrete camphor in containing only half as much oxygen. Its density is less than that of the solid camphor. In composition, the latter differs from oil of turpentine only in the presence of two atoms of oxygen; liquid camphor in the presence of one atom of the same element.

5110. An interesting account of this substance will be

found in the United States Dispensatory.

5111. Other volatile oils, besides that of the camphor tree, yield stearopten analogous to camphor. Of such oils Kane gives the following table:—

when I used a mixture of six parts of alcohol, and one of oil of turpentine, in an Argand lamp.

Subsequently, however, on being consulted, I objected to the use of the continuous on account of the danger arising from its liability to inflame. Experience is shown, by many melancholy disasters, that this counsel was correct.

| nt giving the Camphor.     | Sp. gr.<br>as Liquid. | Melting<br>Point.                 | Boiling<br>Point.            | Sp. gr. of Vapour. | Formula.                                                                                                                                                                                |
|----------------------------|-----------------------|-----------------------------------|------------------------------|--------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| e (Otto) . sley Florentina |                       | 77°<br>70°                        | 550°<br>552°                 |                    | CH<br>C19 H7 O4<br>C4 H4 O                                                                                                                                                              |
| campane                    | 1.014                 | 108°<br>104°<br>68°<br>64°<br>91° | 530°<br>428°<br>430°<br>406° | 5680<br>5455       | C <sub>20</sub> H <sub>10</sub> O <sub>2</sub><br>C <sub>20</sub> H <sub>10</sub> O <sub>3</sub><br>C <sub>16</sub> H <sub>11</sub> O <sub>4</sub><br>C <sub>16</sub> H <sub>14</sub> O |
| rpentine .                 | 1.057                 | 302°                              | 811°                         |                    | C <sub>20</sub> H <sub>20</sub> O <sub>4</sub> C <sub>19</sub> H <sub>14</sub> O                                                                                                        |

mphors arise from very various causes; in some cases they are isomeric with is, in others oxides of them, and in others hydrates; thus, the camphor of turne may be formed at will, by agitating the oil with water and then exposing it d; the hydrate crystallizes out in colourless prisms, sometimes of great size. he peppermint camphor has been found to yield, by the action of reagents, a of compounds. Thus, by the action of glacial phosphoric acid, or of oil of, a light oil was obtained, having the formula C21 H18, which is termed meaby the action of chlorine, a thick heavy liquid is produced, C21 H14 C16 O3. tric acid, menthen yields a heavy oily liquid, C21 H18 O9, which possesses acid ties; and with chlorine, menthen yields a syrupy yellow liquid, having the la C21 H13 C15."

## Artificial Camphor.

112. If one hundred parts of oil of turpentine, refriged by snow and salt, be saturated with chlorohydric gas, by means of an impregnating apparatus, a quantof the gas, equal to about one-third of the weight of turpentine, is absorbed. Meanwhile the turpentine is nged into a soft crystalline mass, from which, allowing drip for some days, about twenty parts of a colouracid liquor are obtained, charged with many crystals, one hundred parts of a white, granular, crystalline subce, which so much resembles camphor in odour and tility, that it has received the same appellation.

redden litmus. It may be sublimed, but not without tial decomposition. If passed through an incandescent e, it is resolved into its constituents. It dissolves in alol, and is precipitated from it by water unchanged. orine is disengaged from it by nitric acid. This subser has been analyzed both by Dumas and Oppermann. ording to the former chemist, it is composed of one me of chlorohydric acid united to one volume of a

compound, formed of ten atoms of carbon and eight of hydrogen, and consequently identical in composition with pure oil of turpentine.

#### Of Camphene or Camphelene, and Terebene.

5114. From artificial camphor, by subjection to the distillatory process with quick-lime, an oil separates, called, by Dumas, camphene, by others, camphelene. This oil is identical in composition with pure oil of terpentine, and differs from it so little in properties, that were it not that the latter has a power of causing a pencil of polarized rays to turn to the kal, of which power the former is devoid, one could not be distinguished from the The liquid from which the artificial camphor crystallizes, has the smell of camphor no less than the crystalline portion, and consists of nearly the same ultimate elements, united to chlorohydric acid. It has a relation to artificial camphor like that which the eleapten of a volatile oil bears the stearopten. When this liquid artificial camphor is distilled with selphuric acid at as low a heat as possible, an oil is obtained, called terebese, which is, like camphene, devoid of the power of causing any rotation in polarized rays. Yet either terebene or camphene, by uniting again with chorohydric acid, may regenerate each the kind of artificial camphor from which it was evolved, and in this respect they differ from each other, which differing from the pure native oil as already stated. Yet, by combining with chlorine, both camphene and terebene acquire a power of causing in polarized light, a rotation in a direction the opposite of that produced by native oil of turpentine (4052).

## Of Kreosote.

5115. This name has been given to an essential oil, to which allusion has been above made, as one of the products of the destructive distillation of vegetable matter. It is represented as highly interesting and important, on account of its efficacy as a medicine, and in preserving meat; being in fact considered as the principle to which pyroligneous acid and wood smoke are indebted for their antiseptic powers, and tar-water for its medicinal virtues.\*

5116. Kreosote is elaborated either from crude pyroligneous acid, or from wood tar, by a series of distillations,

and subjection to different agents.

5117. Besides its activity in medicine, kreosote is alleged to have energetic powers as a chemical agent. It is an oleaginous, colourless, transparent, and highly refracting liquid. It has the smell of crude pyroligneous acid, or of smoked meat, and its taste is caustic and burning. To

The antiseptic power of oil of cloves, and still more that of oil of cinnamon, see equal to those of kreosote, agreeably to my experiments made with meat or cress. A few drops of cinnamon oil added to a paste of gum tragacanth, will prevent, is months, the fetor which otherwise is soon acquired.

that of the oil of almonds. It is rather heavier than ater, being of the specific gravity of 1.037. It boils at 97°.

5118. Kreosote is devoid of acid or alkaline reaction. Vith water it forms two combinations—one a solution of ne part of kreosote in four hundred of water, the other a dution of one part of water in ten of kreosote. It unites all proportions with alcohol, ether, and naphtha, and is apable of dissolving a large quantity of iodine and phosporus, and likewise sulphur, especially when assisted by eat. Agreeably to Thenard, the composition of kreosote expressed by the formula, C<sup>14</sup> H<sup>9</sup> O<sup>2</sup>.

## Of Essential Oils which are Hydrurets.

5119. Among the oils which may be called hydrurets, re the hydruret of benzule or oil of bitter almonds; an ily hydruret existing in the commercial oil of cinnamon r cassia, called hydruret of cinnamyl; the oil of spirea lmaria or hydruret of salycyl; and the hydruret of cumyl, erived from the oil of cumin. Of the three first menioned oils, some account has been given in treating of heir radicals; and to the hydruret of cumyl, allusion was nade in paragraph 5103. I do not, however, deem it exedient to give any details here respecting any of these ils, excepting the hydruret of benzule. Of this I shall reat for the purpose of exemplification.

## Of the Hydruret of Benzule, or Oil of Bitter Almonds.

5120. The formula of this hydruret is C<sup>14</sup> H<sup>5</sup> O<sup>2</sup> + H, or BZ + H. By listilling bitter almonds, or the leaves of cherry laurel, with water, a volatile product comes over, consisting of a mixture of the hydruret of benzule, of benzoic acid, of gum benzoin, and cyanhydric acid. In order to extrinte the hydruret from this mixture, a second distillation is requisite, with the previous addition of chloride of iron, hydrate of lime, and sufficient water o liquefy the whole. Under these circumstances, the oil may be distilled, accompanied by water, which may be separated by the usual means, and subsequent agitation with chloride of calcium.

5121. Properties.—This hydruret is colourless and transparent, refracting light strongly, being endowed with a strong odour like that of cyanhydric acid, and a hot taste. Its specific gravity is 1.043; its boiling point 356°. It is soluble in thirty parts of water, and in alcohol in proportion. Its vapour may be transmitted through a red-hot tube without decomposition. It burns with a white, though smoky flame. By absorbing two steems of atmospheric oxygen, one to unite with an atom of hydrogen, the other to take its place, this hydruret is converted into benzoic acid. Sub-

jected, at a high temperature in close vessels, to hydrate of potash, it forms a benzoate of that base by absorbing the oxygen, and liberating the hydrogen of an atom of water.

5122. The hydruret of benzule undergoes no change by being in council with aqueous solutions of caustic alkalies or earths, but, while thus situated, a few drops of cyanhydric acid will enable crystals of benzoin to be general.

rated.

- 5123. By contact with chlorine or bromine, the hydruret of bearule is converted into chloride or bromide, its hydrogen being simultaneously converted into chlorohydric or bromohydric acid, by uniting with one or the other of those elements.
- 5124. An iodide of benzule can be obtained by the reaction of the distribution of this compound radical with the iodide of potassium; in like master a sulphide, by the distillatory reaction of a chloride with the sulphide of lead; and a cyanide, by substituting a cyanide of mercury and resorting to the same means.

#### Of the Amiduret of Benzule or Benzamide, BZ NH.

5125. From the preceding formula it must be evident that the composed of which the name is above given, consists of benzule, and the composed radical, amide.

5126. This amiduret arises from the reaction of the chlorides of the reaction 
puric acid with the peroxide of lead.

5127. Amiduret of benzule crystallizes in right rhomboidal pearly prime or tables. A hot concentrated solution by refrigeration, yields a soft mest of very fine crystalline needles, which are gradually transformed into broad colourless laminæ. These crystals melt at 239° into a colourless lique, and at higher temperatures are susceptible of forming an inflammable very

pour. They are soluble either in water, alcohol, or ether.

5128. Water being present, alkalies or acids resolve this amiduret into ammonia and benzoic acid. On being heated with anhydrous baryta, a benzoate of this base is produced, with a disengagement of ammonia, much heat, and the volatile oil called benzole. Similarly treated with potassium a cyanide of this metal results, with the evolution of an oleaginous aromate liquid of a slightly sweet taste. The hydruret of benzule unites also with anhydrous formic acid, generating a compound acid called formobenzole acid; also with benzoic acid, forming what is by Liebig termed a benzoale of the hydruret of benzule.

## Of Resins.

5129. Resin is now the generic name of a class of bodies, of which common resin or rosin is an exemplification, having had its name extended to the class in consequence of their analogy with it. On this account, English writers have latterly used the word resin, generally employing the word rosin as the name for the substance which formerly was designated either as resin or rosin. In pharmacy, rosin is also known as colophony or colophonium; especially on the continent of Europe.

5130. Resins are found in vegetables and in the fossil tate, as in the instance of amber and asphaltum; but in very case, are considered as having been originally the

products of vegetation.

5131. In vegetables, resins exist more or less in combination with essential oils; and I believe them to be geneally produced by the reaction of such oils with oxygen. It has been mentioned that, when distilled per se, almost esinous substance, which does not come over. It is also rue, that any resin, exposed to destructive distillation, gives ise to more or less pyrogene oils of the volatile kind, as well as carburetted hydrogen, and carbonaceous depositions, and residues.

5132. In many cases, as in that of the turpentine of commerce, the compound formed by the resin and the rolatile oil with which it is naturally associated, is sufficiently liquid to flow from incisions made through the mrk and sap wood. It is thus that the copious supply of urpentine found in commerce, is obtained from the long-

eaved pine of the Carolinas.

5133. Another portion of resinous matter, expelled by ire, forms the tar of commerce. This contains some renarkable volatile compounds generated by heat, called paraffine, cupion, and kreosote. The former is a concrete oil, the others liquid. Tar also contains acetic acid in combination with the several peculiar resins, called pyre-

ene, or pyrogene resins, by Berzelius.

5134. As the expulsion of resinous matter by the tarproducing process destroys the peculiar properties of resins, I believe it is not resorted to in obtaining resins in other cases. More valuable resins, which do not spontateously exude, are generally extracted by digesting the regetable product containing them in alcohol. From the alcoholic solution, when it takes up other substances, the resin is precipitated by water.\*

The celebrated varnish of the island of Japan exudes from the rhus vernix, which is among the forest trees of the United States, being notorious for its poisoness influence on some persons, while to others comparatively harmless. The active rinciple to which its poisoning power is due, would be a worthy object of investigation by any one not susceptible of the injurious effects alluded to. In the art of jamaning in this country and in Europe, other substances are made to imitate the first of the real Japan varnish, named from the country in which it is employed.

5135. Resins are all insoluble in water, and for the most part, directly or indirectly, soluble in alcohol, and in volutile and fixed oils. They cannot, like volatile oils, be detilled with the aid of water. When subjected, pur se, to the distillatory process, they are decomposed, as above mentioned, into carburetted hydrogen, carbon, peculiar resins, and volatile oils, some acids, and more or less carbon partly in the state of lamp-black, partly in union with the other products, whence their dark or black colour.

5136. In few instances do resins assume a crystaline form. They are brittle when pure, and generally transferent, rarely colourless, having, commonly, various hues of yellow or brown, but sometimes green or red. There is a great resemblance in properties between resins and con-

crete oils, such as suet, tallow, spermaceti.

5137. Resins are distinguished by a greater hardness and tenacity, and in being sticky to the touch instead of being greasy. Hence rosin serves to create the necessary attrition between the hair of the bow and the strings of the violin, which is an effect the opposite of that for which oil is used in machinery. In this, as well as in other respects, wax approaches the resins in character more than any other concrete fixed oil. But this adhesiveness is much increased by heat, so that at ordinary temperatures copal, amber, and many other resins, are not sticky. In consistency resins much resemble gums, but are distinguished from them by insolubility in water, and solubility in fixed and volatile oils, and generally in alcohol and ether.

5138. Some resins resemble fixed oils, in containing two substances, of which one is more soluble, the other less soluble in alcohol. This characteristic is, in some instances displayed in their habitudes with some essential oils. Besin, for instance, is said to be only partially soluble in

naphtha.

5139. Resins, also, are susceptible of saponification. of far as to combine with alkaline and other bases formed salts, in which the base, being imperfectly neutralized, possesses the detersive power. It is well known that rosin is a constituent of common brown soap, yet, according to Ure, it cannot enter into it advantageously beyond the proportion of a third. There is this important difference, however, in the phenomena of the reaction of fixed oils

bases, and that of resins, that there is no base to be

led analogous to the oxide of glyceryl.

- 40. Concentrated nitric acid and resins react, in some 3, with an explosive ignition. According to Berzelius, dissolve in concentrated sulphuric acid, when cold, out decomposition, although when hot reciprocal deposition ensues. I have ascertained that sulphuric forms, either with oil of sassafras, or with oil of s, resins, by which it is coloured to a miraculous desince a six-millionth part suffices to create a rosy A similar effect, in an inferior degree, ensues from resence of oil of cloves. To the resins thus produced, re given the names of sassarubrin and cinnarubrin. lieve in any case it will be found, that more or less is produced by the reaction of concentrated sulphuric with essential oils. In fact, such oils, to a certain it, act as bases to this acid, diminishing the sourness diluted solution, and when such a solution is saturated ammonia, a resin formed from the oil separates.
- 41. Resins are soluble without alteration, either in c or chlorohydric acid.\*

42. Prof. F. W. Johnson has proposed to represent sins by two general formulæ, either of which contains

appears from Unverderben's laborious investigations, that by the various use or hot alcohol or ether, resins, as they are found in nature, may be resolved rious substances, differing from each other as respects readiness to combine uses; so that he has classed them as resins strongly electro-negative, modeslectro-negative, feebly electro-negative, and indifferent. This author founds versity of designation, on their greater or less disposition to combine with ua, carbonate of soda, or caustic alkaline solutions.

eably to Johnson's Report to the British Association, for 1832, Buchner and ger had described some resins as having weak basic properties. Resins exfrom jalap and euphorbium had each been found a compound of two resins s acid, the other a weak base: also all drastic gum resins were considered by

hemists as similarly compounded. well known that all resins are electrics, and by friction become negatively

ied. ording to the author last mentioned, sandarach is a mixture of three resins;

five; benzoin of three; guiac of two; and lac and colophony of several. n rosin or colophony is subjected to cold alcohol, of the density of 867°, one dissolves, called alpha resin or pinic acid; while another remains, called beta r sylvic acid. By exposing pinic acid to distillation, another acid is generated colopholic. Again, the solution of pinic acid may be decomposed by acetate er, of which the oxide precipitates with the acid, leaving an indifferent resin tion.

white rosin, from the pinus maritina, consists of an acid, crystallizable resin, pimaric acid. Distilled in vacuo, pimaric acid gives rise to another, called wie acid. Boiled with nitric acid, pimaric yields azomaric acid. But there is to the variety of compounds resulting from subjecting resins to heat and vaolvents. It may be of some practical importance to know, that resins are not encous substances, and that even the rosin of different trees may contain difncide.

forty atoms of carbon, while one holds from sixty to sixtyeight atoms of hydrogen, with from one to twenty of oxygen; the other, forty to fifty-four of hydrogen with from

seven to fourteen atoms of oxygen.
5143. He infers, that the resin of scammony, C. H. O., extracted from crude scammony by alcohol, contains the largest quantity of oxygen of any resin hitherto analysed; and that the resin of jalap, obtained by evaporating the alcoholic extract, and subsequent boiling in water, of which the formula is C<sup>40</sup> H<sup>34</sup> O<sup>18</sup>, is, as respects the quantity of contained oxygen, surpassed only by scammony.

5144. Agreeably to the same author, there is a striking relation between the formulæ of the resins of ammoniae and asafætida, the former being C40 H25 O9, the latter, C

H<sup>26</sup> O<sup>10</sup>, as if the one were merely a hydrate of the other.
5145. Berzelius considers our knowledge of the composition of resins as yet too imperfect to justify us in placing much confidence in these suggestions of Johnson as to the grouping of all resins under two formulæ as above mentioned. Report for 1841, 171.

5146. The following list of the more important resins of commerce, with their formulæ, is taken from Kane's Ele-

ments, p. 969.

| , P               |                   |                   |                          |
|-------------------|-------------------|-------------------|--------------------------|
| Anime Resin       | C40Hs2O           | B. Sandarach      | CooHarOs                 |
| Elemi Resin       | •                 | A. Euphorbium     | } C-n-o                  |
| Fossil Copal      | C40H32O           | <b>Asphaltene</b> | CeoHraOe                 |
| B.* Mastic Resin  | C49H31()3         | A. Olibanum       |                          |
| Antiar Resin      | C40H30O3          | Labdanum          | CeHaO:                   |
| B. Copal Resin    | C40H31O3          | Pasto Resin       | C+oHzaCh                 |
| Birch Resin       | C40H33O3          | Sagapenum         | (.mHad),                 |
| A. Mastic Resin   | C40H31O4          | Scammony          | CAOH 200m                |
| Copaiva Resin     | Cash of           | Jalap Resin       | Crell 200 la             |
| A. Elemi Resin    | C40H32O4          | Galbanum          | (.4.Hr.G.                |
| B. Olibanum Resin | •                 | Dragon's Blood    | CoHath                   |
| C. Sandarach      | C40[130()8        | Gamboge           | C•oH•O•                  |
| Ammoniac Resin    | C40H24()9         | A. Asafortida     | C+oHa()ra                |
| B. Asafætida      | C40H28()9         | Acaroid Resin     | C.⊷Ha.On                 |
| Guiacum           | C45H93()10        | Opoponax          | ('**'H**0'*              |
| Bdellium Resin    | } C40   I 21 () 2 | B. Benzoin Resin  | (_^4\[]as() <sub>0</sub> |
| A. Sandarach      | 3 Callanda        | A. Benzoin Resin  | C4vH=0                   |

## Of Wax.

5147. This word is generally used to designate the substance of which bees make their honeycomb; more accu-

<sup>\*</sup> Where a native resin has been separated into two, by solvents, the letters A and B are used to distinguish one from the other.

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es of the leaves of certain trees, and the skins of certain ne fruit; also to be yielded by the cabbage, and in a ge proportion by the berries of several species of the rtle, myrica angustifolia, latifolia, and cerifera.

pollen of flowers swallowed and excreted by bees; but as been proven that the wax of bees is secreted by an situated on the sides of the medial line of the abdo-

as been proven that the wax of bees is secreted by an an situated on the sides of the medial line of the abdon of the insect. On raising the lower segments of the lomen these sacs were observed; also the scales, or ngles of wax arranged in pairs upon each segment.

. Huber ascertained that bees, while prevented from ng abroad in quest of food, and fed solely on sugar,

re capable of generating wax.

that myrtle wax yields, by saponification, stearic, rgaric, and oleic acids, and glycerine, like a true fat, ile, when subjected to the same reagents, bees-wax is able only of a partial saponification, yielding in lieu of congener, of ethal, or of the sweet principle of oils, a stance called cerain, which differs neither in composin nor properties from that portion of wax which is inso-

le in boiling alcohol.

tion dissolved in the hot alcohol is called cerine. It is ine only, that is capable even of the partial saponificanto which allusion has been made. As respects this saration into cerine, and myricine, by boiling alcohol, sowax resembles a fat, consisting of stearine and marine, while devoid of oleine; but in its chemical constituant habitudes, with bases, it resembles the resins. ax is also destitute of the greasiness or slipperiness of tending, when interposed between surfaces, to impede ir sliding, rather than to facilitate it like an oil. Upon whole I consider bees-wax as a substance intermediate tween a concrete fixed oil and a resin.

5151. The yellow wax of commerce is obtained by fug, and washing, the crude wax of the comb with boiling ter. Yellow wax is converted into white wax by causing o form thin ribbons by flowing while melted upon a reving wooden cylinder, half immersed in water, and sub-

sequently exposing these ribbons to the solar light and the air, as in the old process for bleaching linen. The war of the honeycomb, before being supplied with honey, is white.

5152. Pure white wax is of the specific gravity 960, 966. It is insipid and inodorous, insoluble in water, partially soluble in boiling alcohol, and perfectly soluble in essential or fixed oils. It fuses at about 154°. Its general uses are too well known to need description. Not being much acted on by acids, it is used to defend corks, and as ceneral or lute, for chemical apparatus.\*

#### Of Caoutchouc or Gum Elastic, and Caoutchoucine.

made in certain trees, and congeals in the form of the mould upon which may be received. Like essential oils, devoid of oxygen, it consists only a carbon and hydrogen, Cs H7. As respects its chemical habitudes, it might be considered as a resin, were it not for its wonderful and peculiar elasticity, and the mechanico-chemical property of allowing gases to get through its pores with a celerity not corresponding with the minuteness of the atomic weights. In its native state, instead of being held in solution, as resins are usually, by an essential oil, it is merely suspended in water, as the ter and caseine are in milk. Faraday found in a portion of caoutchow milk which he examined, the following ingredients:—

| wn bitter azotized<br>precipitable by ni          | • |                 | • | • |   | 7.1  |
|---------------------------------------------------|---|-----------------|---|---|---|------|
| ctable albumen,                                   | • | · <b>7</b><br>- | • | • | • | 1.9  |
| Substance soluble in water, insoluble in alcohol, |   |                 |   | - | • | 2.9  |
| Water holding a small quantity of free acid,      |   |                 |   | • | • | 56.3 |
| Caoutchouc, -                                     | • | •               | • | • | • | 31.7 |

5154. Pure caoutchouc, carefully prepared from the native emulsion. • of the density of .925, being transparent and colourless, and, when in mess yellowish white.

5155. It is utterly insoluble in water or alcohol, but soluble in pure effer (oxide of ethyl), and likewise generally in pure essential oils, especially oil

"Of Cerosic.—Mr. Avequin has examined the wax which covers the sugar cane, and the lower part of the leaves by which it is surrounded. It may be obtained by scraping the surfaces covered with it. In the violet variety of the plant in questions this wax is so abundant, that it is inferred by Mr. Avequin that it might be profitably collected for the purpose of making candles. The scrapings are digested in colder cohol to remove impurities. Afterwards they are dissolved in boiling alcohol. The solvent being removed by distillation, the wax is isolated.

This wax is slightly yellow, hard, brittle, easily reducible to powder of a brittle, fuses at 176°, and burns like ordinary wax or spermaceti. It is less solubies ether than alcohol. From boiling solutions in either solvent, it separates in performedle-shaped crystals by refrigeration. Mr. Avequin proposes for this wax is name cerosie, from the Greek ceros, wax. The formula of this wax is, according to

analysis by Dumas, C<sup>20</sup> H<sup>30</sup> O<sup>1</sup>.

sassafras, cajeput, and turpentine. It does not, however, readily liquefy, t, absorbing many times its bulk of the solvent, may be liquefied aftertrds by rubbing through a sieve. It is, perhaps, even more soluble in the rogene oils, such as naphtha, whether native or as obtained from coal, d in the peculiarly volatile oil, called caoutchoucine, which it yields itself r destructive distillation, and repeated subsequent rectifications. It has en mentioned, that this oil was lighter than any analogous native product. is, in fact, lighter and more volatile than common ether, its density being dy 670, and for its boiling point 90°. From none of the volatile oils, not ren caoutchoucine, have I recovered caoutchouc, without more or less derioration. This may be presumed to arise from a minute quantity of renous matter formed at the expense of the solvent which remains with the I have found a great diversity in the solubility of caoutchouc. either in caoutchoucine, nor in ether, have I found the ordinary bag caoutnouc to dissolve readily. It softens and swells up, but does not liquefy. ut a large lump of massive caoutchouc, sent to me from London by Mr. inderby, was readily liquefied either by the one or the other of the last mentioned solvents, and by the ether was deposited in a perfect state. enve not learned the source of the more soluble caoutchouc thus alluded to, or have I met with any notice respecting this difference of solubility. **aoutchouc** burns with an excessively fuliginous flame in atmospheric air, text in oxygen gives an intense light by the oxidation of the carbon forming be smoke (645). When fused, per se, it is converted into a tarry matter, thich does not indurate by drying. This tar may be ignited by nitroso-Mric acid.

5156. Dr. Mitchell ascertained that caoutchouc bags, after soaking in a mixture of ether and alcohol of the specific gravity of from 750° to 780°, or the usual officinal strength, may be inflated with air, and the material of which they consist consequently extended to various degrees of tenuity, actording to the peculiar character of the variety subjected to trial. Hence may be used to make balloons, gas bags, or sheet gum elastic, which is vary useful for fillets, with which to make air-tight junctures or lutings. There is no better mode of joining a tube to the tubulure of a retort, or reviver, than by tying about the tubulure the body of a small caoutchouc ag, while the tube is inserted into the neck, and carefully secured by a gature. Fused caoutchouc is useful in some cases as a lute. It will not, nowever, resist fuming nitroso-nitric acid.

5157. Dr. Mitchell has made some very interesting observations respectag the power of gases to pass through thin membranes of caoutchouc.
by some inconceivable process, gases, which are all prone, in a greater or
as degree, to reciprocal intermixture, will effect this result, notwithstandby the interposition of caoutchouc, and the opponent influence of great

ressure.

losed by a caoutchouc membrane, was introduced into a vessel of hydromen, this gas made its way into the vessel, until the membrane burst outpards; but when the vessel, while similarly closed by the membrane, and eplete with hydrogen, was exposed to common air, the hydrogen escaped stil the membrane burst inwards. A tube, with a trumpet-shaped mouth, sing bent so as to form a syphon, and the larger orifice closed by the membrane while full of atmospheric air, a suitable quantity of mercury was bursed into the syphon, until it stood in both legs at the same height. Unter these circumstances, when the membrane was brought into contact suc-

cessively with different gases, they were found to enter with various degree of celerity, as will appear from the following statement:—

|                     |    |   |   |   | H. | X.             |
|---------------------|----|---|---|---|----|----------------|
| Ammoniacal gas      | •  | • | • | • | 0  | 1              |
| Sulphydric acid     | •  | • | • | • | 0  | 21             |
| Cyanogen -          | •  | • | • | • | 0  | 31             |
| Carbonic acid       | •  | • | • | • | 0  | $5\frac{3}{2}$ |
| Protoxide of nitrog | en | • | • | • | 0  | 61             |
| Arseniuretted hydi  |    | • | • | • | 0  | 27             |
| Olefiant gas        | -  | • | • | • | 0  | 28             |
| Hydrogen -          | •  | • | • | • | 0  | 371            |
| Oxygen -            | •  | • | • | • | 1  | 13             |
| Carbonic oxide      | •  | • | • | • | 2  | 40             |
| Nitrogen -          | •  | • | • | • | 3  | 15             |

5159. The gases continued in some instances to enter until the mescay

in the longer leg rose to the height of sixty inches-

5160. It is quite surprising that the atoms of ammonia should put through the membrane with greater celerity than those of hydrogen, who each of the former consists of three of the last mentioned gas, united with one atom of nitrogen. Also that two atoms of oxygen, while associated with an atom of carbon, should permeate the membrane more speedily than isolated atom of oxygen.

5161. It also appears from experiments made by Dr. Mitchell, and peated by myself, that caoutchouc is probably more highly susceptible electric excitement, than any other organized body; and probably is at least

equal in excitability to any inorganic substance.

## Of Balsams.

- 5162. The word balsam has been used to designate metive solutions of resinous matter in essential oils, which, like the turpentine of commerce, exude spontaneously from trees or shrubs.
- 5163. Among these, however, there are some distinguished by the presence of benzoic or cinnamic acid, or both. It is to the balsam of Peru and Tolu that this remark applies particularly (3060). Styrax has also been alleged to contain a minute proportion of benzoic acid, but is not included among balsams by Soubieran, and by this author the corresponding French word baume is employed to designate artificial compounds of resins with an acid and volatile oil.
- 5164. According to Fremy, balsam of Peru consists of resinous matter, of cinnamic acid, a liquid essential of called cinnameine, and a crystallizable oil supposed to a hydrate of cinnamyle (3058), called metacinnameine.

5165. Balsam of Tolu consists of resin, cinnameire

cinnamic acid, and, perhaps, metacinnameine.

5166. Balsam of copaiva, or copaiva balsam, consists

of a volatile oil, and two resins without any acid.

5167. But agreeably to the investigations of Deville, benzoic acid also exists in the two first mentioned balsams, and when the balsam of Tolu is distilled, per se over a naked fire, a volatile oil, and likewise benzoic ether, are obtained. It is suggested that the resin of the balsam is an oxide of this ether.

5168. It appears that, by reaction with caustic potash, cinnameine is resolved into cinnamic and benzoic acid in union with the alkali: an oily substance, little soluble in water, called peruvine, being simultaneously evolved. There is some analogy between this process, with its re-

sults, and those of saponification.

5169. By some authors the word balsam is restricted to resiniferous liquids containing benzoic acid. It might be more reasonable to consider an acid of some kind as requisite, yet it is evident that ordinary acceptation does not justify the idea that the presence of an acid is necessary.

## Of Gum-resins.

5170. This name is applied to a class of vegetable substances, which consist of a mixture of resin, gum, essential oil, and extractive matter. Opium, aloes, ammoniac, asafatida, euphorbium, galbanum, gamboge, myrrh, and scammony come under this head.

5171. As the resin and essential oil require alcohol, the gum and extractive matter water, for solution, proof spirit

is the best solvent of the gum-resins.

## Of Opium.

5172. This complex substance contains the following proximate principles;—

1. Morphia, in the state of neutral sulphate, and super-

meconate.

2. Paramorphia.

3. Pseudomorphia.

- 4. Codeia, in the state of supermeconate.
- 5. Narcotina.
- 6. Narceia.
- 7. Meconin.
- 8. Meconic acid, partly combined with bases.

- 9. Ulmin.
- 10. A peculiar resin.
- 11. A fatty oil.
- 12. Caoutchouc.
- 13. Gum.
- 14. Bassorin.
- 15. Lignin.
- 16. The sulphate of potash, lime, and magnesia.
- 5173. Of these substances, morphia, paramorphia, pendomorphia, codeia, narcotina and narceia are ranked as vegetable alkalies, all having the power of neutralizing acide. Meconin is an indifferent or neutral subtance, which was announced to exist in opium in 1832, by M. Couerbe, but which is found to be identical with the crystallizable principle of M. Dublanc, jun., discovered several years before. Paramorphia, pseudomorphia, narceia, and meconin, exist in opium in very small amount. For a method of detecting opium, see meconic acid (5265).

# Of Bitumen, Petroleum, Naphtha, Amber, and Mineral Coal.

- 5174. There is in nature a gradation of substances, apparently arising from the wreck of a former world, from naphtha, which is highly volatile, to anthracite, which is extremely insusceptible of the aëriform state. Possibly the diamond may be considered as terminating the series; as it has been suggested to result from the decomposition of vegetable matter.
- 5175. Bitumen, in a concrete state, is exemplified by asphaltum. The coal called bituminous, owes to the presence of bitumen its capability of caking, and yielding carburetted hydrogen when ignited. Bitumen is found also in a tarry state, or more or less liquid, according to the quantity of petroleum with which it may be united. Caking coal may be considered as a compound of carbon with bitumen, and a minute portion of silex and iron, and sulphur: anthracite, as consisting of the same ingredients, substituting water for bitumen, though in a lesser proportion.
- 5176. Petroleum, or naphtha, is the name given to an inflammable liquid which rises out of the earth like spring water, so that some wells cannot be freed from it. The name of naphtha is more properly given to a very volutile oil which may be obtained from petroleum by cautious distillation, preferably with water. Besides more or less bitumen, by which it is discoloured to a greater or less degree; agreeably to the researches of Pelletier and Walter, petroleum comprises three volatile oils, and a species of paraffin. The names, boiling points, and formulæ of the oils, are as follows:—naphthel, C. H., boils at 384°; naphthene, C. H., boils between 185° and 194°.

of acids. 453

5177. Naphtha proper.—The last mentioned oil may be considered as the true naphtha, being the liquid employed for the preservation of the metals of the alkalies. It much resembles oil of turpentine in properties and composition. Potassium, of which the specific gravity is .865, sinks readily in naphtha.

5178. During the destructive distillation of bituminous coal, a bituminous liquid, called coal tar, condenses, from which an artificial naphtha may be

extricated, which is used as a solvent of caoutchouc.

5179. Seneca Oil, American Oil.—Under these names two liquids are now to be met with in commerce. The former is obtained from the vicinity of the lake after which it is named; the latter from a well in Kentucky, which was sunk for the purpose of obtaining spring water. Either yield, by distillation with water, more or less naphtha, and contain heavier oils

requiring a higher heat to bring them over by distillation.

vegetable matter. It is distinguished by burning with a peculiar odour, and yielding, when subjected to distillation, succinic acid, and a peculiar essential oil, called oil of amber, which resembles crude naphtha in smell and other properties. The acid sublimes into the neck of the retort in crystals. Amber is insoluble both in water and alcohol. Dr. Kane suggests that it may be the turpentine of an extinct species of tree, belonging to a former seological epoch. It would seem rather to be a variety of copal, which it much resembles in appearance and properties, as that the one may be mistaken for the other, on superficial examination.

#### OF ACIDS.

## Of Acids relatively to the Proportions of Base required for their Saturation.

5181. It has long been known, that certain acids, such for instance as nitric, or chloric acid, cannot be isolated so as neither to be in unison with water, nor with any other oxide acting as a base. Until of late, however, it does not seem to have been perceived, that the water in such acids must act as a base. Now it is held, that wherever water, unless replaced by another oxide, cannot be expelled from n acid without a decomposition of the acid, or a destruction of its properties, such water, while combined with the acid, must be considered as acting as a base. Moreover, as when one atom of water, or other oxide, is found indispensable to the existence of an acid, that one atom has been considered as performing a basic part, so, consistently, when two or three atoms of water or other oxide are ascertained to be no less necessary, the two atoms, or three atoms of water or other oxide thus required, are considered as acting as bases. Experience has shown that in this way, some acids require one, others two, and others

three atoms of base, and are called accordingly monobs

bibasic, or tribasic acids.

5182. But, it may be inquired, how is this diversity: the acids ascertained? The answer is, by ascertaining loss of weight which they sustain, on combining with base to saturation. Of course, the weight of the formed with a dry base, should be the aggregate weight that base and the anhydrous acid. This may be found desiccating the resulting salt. The difference between weight of this saline aggregate, and that of the sam of the weights of the hydrated acid and dry base, must be duth

the escape of basic water.

5183. Although when water, which can be replace another base, is essential to the existence of an some follows that it must be considered as basic; the state ought not to infer that it cannot act as a base to 🖎 which can exist without it. Both sulphuric and phoric acid unite with water as a base, although care of existing in the anhydrous state. This prelimina planation having been given, it is hoped that the dent will be prepared to understand the following ment, respecting the three classes of acids above

5184. Acids, as respects the quantity of base with which they are capable of combining, may be divided into the classes. Those requiring one equivalent of base, called monobasic; those requiring two equivalents, called bibasic; those requiring three equivalents, called tribasic acids Water acts as a base in combining with any acid of either class, and is subject to the same laws as other bases.

5185. The compounds, intherto called hydrated audit are in combination with one, two, or three atoms of but water, accordingly as they belong to the monobasic, and

bibasic or tribasic class.

5186. When the hydrate of an acid of either kind is prosented to a base, capable of displacing water, for every atom of the new base which unites with the acid, an attention of water must be expelled. As the single salts of most basic acids can have only one equivalent of base, so them there can only be one kind of base; but in biball acid salts the equivalents may be of one kind only, or 🥊 two kinds; and in tribasic acid salts, of one kind, or

o, or of three kinds. In either case, water, acting as a se, is liable to be present in the same proportions as any ner base, and may replace or be replaced by other bases. I that has been said of water, is also true in many cases oxide of ammonium.

5187. Different bases, salified by the same monobasic id, may combine to form double salts. Of course, salts ving water for their base are not excepted; but double ts thus formed with an equivalent of basic water, on acunt of their sourness or reaction with litmus, have been led acid salts. When in such salts the water is reced by another base, two neutral salts result, which be separated by crystallization, provided they differ solubility, and crystallize separately, in forms sufficient-different to be distinguished.

5188. When monobasic acids are united to more than equivalent of base, not being neutral, as bibasic or trisic acids are, with the same number of basic equivalents, y are called basic salts; which conveys the idea of a t consisting of an acid united to one or more atoms of se in excess. Yet when the atoms thus situated, are sented to another atom of the same monobasic acid, in state of hydrate, they can displace no more than one m of basic water; for this obvious reason, that there is be no more than one atom of basic water in union

h such an acid.

5189. Salts of bibasic acids, when one of their atoms of se is water, are, from their sourness, called acid salts; t, substituting another base for water, does not produce louble salt. For this, two atoms of acid and four atoms base would be requisite.

5190. Acids produced by dry distillation, are called pyene acids. Such acids are rarely created by subjecting nobasic acids to that process; but pyrogene acids,

en thus created, are always monobasic.

5191. Under like circumstances, bibasic acids give birth en to two new monobasic acids, as in the instance of lic acid.

5192. By the same process, tribasic acid may give rise three equivalents of a monobasic acid, as in the case of muric acid; or they may be resolved into two monobaccids, or a bibasic and a monobasic acid, as may be in the case of meconic acid.

5193. In the following table, taken from Gregory's big, the anhydrous acid is represented by R, the met oxybase by MO, and water by the usual symbol HO.

#### Formula for Monobasic Salts.

General formulæ for the salts of the bibasic acids.

General formulæ for the salts of the tribasic acids.

#### Of Acetic Acid.

5194. Acetic acid is monobasic, being a hydrated to ide of acetyl (3093), as may be seen from its form C' H' O'+HO.

5195. As the cause of the sourness in fermented liquand various products of vegetation, this acid, having

the whole class of acids; though at this time many of the compounds recognised as acids, are devoid of the attriute on which the general name is founded. Acetic acid the only valuable ingredient in vinegar, causing the souress indicated by its name, which differs but little from in aigre, the words expressive of sour wine in French.

5196. This acid occurs in nature in many products of be vegetable and animal organization: as for instance, in be black elder (sumbucus niger); the pleurix dactilifera, nd rhus tiphinus; in sweat, urine, milk, and the fluids of

he stomach.

5197. It had long been observed that the fermented livors containing the most spirit made the strongest vine-Although pure alcohol is not liable to be acidified er se, when diluted with water holding fermentable subtances, it is readily converted into vinegar. For this urpose each atom requires four of oxygen. Two atoms I this element are requisite to remove two of hydrogen, y which ethyl (3069), the radical of the alcohol, is changed nto acetyl (3093), the radical of acetic acid. At the same ime, two atoms of oxygen are required to be added to the me atom previously in union with the ethyl, to make the bree required for acetic acid, which is a trioxide of acetyl. The formula of alcohol is C4 H5O+HO. If to this we add our atoms of oxygen, we have C4 H5 O5+HO, which gives he formula of hydrated acetic acid = C<sup>4</sup> H<sup>3</sup> O<sup>3</sup> HO+2HO n excess.\* See paragraph 3094 and note.

5198. I shall defer the exposition of the phenomena, auses, and circumstances, on which the conversion of vious liquids into vinegar is dependent, until I treat of fermentation. Practically, every body has a general idea of he mode in which wine, cider, or beer, vinegar, is obtained.

5199. Acetic acid is also a product of the destructive listillation of wood. In that case it forms what has been

Liebig alleges that a strong and agreeable vinegar may be made by exposing to be air for some weeks in a warm situation, the following mixture;—100 parts pater, 13 brandy, 4 parts honey, and I crude tartar. Of course, one part cream of testar might be substituted for the crude tartar.

The scatification of mixtures of vegetable juices with spirit, has been very much impedited, of late years, by a high temperature, and allowing the liquor to drop from table through holes like those of a colander, on beach wood shavings. Respecting in, and other processes for the generation of acetic acid, much information will be in the liquor to drop from the through holes like those of a colander, on beach wood shavings. Respecting in, and other processes for the generation of acetic acid, much information will be in the liquor to drop from the through holes like those of a colander, on beach wood shavings. Respecting in the liquor to drop from the through holes like those of a colander, on beach wood shavings. Respecting in the liquor to drop from the through holes like those of a colander, on beach wood shavings. Respecting in the liquor to drop from th

called pyroligneous acid, which contains various other substances. From these the acid is extricated by combining it with a base, and subsequent distillatory decomposition of the resulting salt by sulphuric acid, the impure acetate having been first cautiously fused to get rid of impurities.

5200. The acetic acid, thus obtained, is much diluted with water, from which it may be freed by digestion with anbydrous sulphate of soda, and subsequent distillation. In this way, according to Liebig, a sufficient degree of concentration may be attained to render the acid crystallizable. As in the case of other organic acids, that in question cancel exist excepting in combination with basic water, or some other base.

5201. The distillation of dry acetate of copper, has been long made the means of evolving the contained acid, in a concentrated state. Resort has also been had to the composition of the dry acetate of soda, or lead, with equivalent portions of concentrated sulphuric acid. 1. According to Liebig, the proportions should be, 3 acetate of soda,

with 9.7 acid: or, 3 acetate of lead, with 8 acid.

5202. Pure hydrated acetic acid crystallizes in shining transparent lamellar, or tabular, crystals. At the temperature of 63° nearly, these crystals fuse into a limpid liquid of the density of 1.063; of which the pungent and distinguishing smell and taste may be inferred, from the inferior effect of strong vinegar. In its concentrated form, as it is capable of blistering the skin, its action upon the tongue must be insupportable. Like other liquids greedy of water, it produces fumes on contact with the aqueous vapour of the atmosphere. It boils at 142°, and unites in all proportions with water, alcohol, ether, many essential oils, camphor, and some resins. When in the state of vapour, it is capable of burning with a blue flame, and being resolved into water and carbonic acid.

5203. It has been mentioned, that, when liquid, crystallizable acetic acid is denser than water. To a certain extent, by admixture with this liquid, a condensation ensues but a further addition of water causes the opposite change. Equal parts by weight have the same density as the pure hydrate. The highest density attainable is 107, indicating the presence of three atoms of water, and one of anhydrous acid; or by weight, 772 acid, and 228 water.

5204. Allusion has been made to the process by which platinum black causes the acetification of alcohol\* (1607).

5205. Of Pyroligneous Acid. The process by which charcoal is obtained by the destructive distillation of wood, has been mentioned as one by which acetic acid is generated. Thus produced, it is generally known as pyroligneous acid, being very much disguised by impurities. In fact, pyroligneous acid so called, contains beside the scetic acid, paraffine, eupione, kreosote, and the pyrogene, resinous matter, called pyretene by Berzelius.

5206. When the process is performed with a suitable apparatus, this acid is collected. Pyroligneous acid may be considered as the matter of wood smoke in the liquid form; and when applied in this state to salted meat, is at least as efficacious as when employed as smoke in the usual way. The process of the smoke-house is less susceptible of precision, and is liable to produce an injurious

rise of temperature.

5207. Of the Acetates.—These salts are soluble, with very few exceptions. Only two are cited as insoluble by Liebig; those of molybdenum and tungsten. The acetates of silver and of the protoxide of mercury, are soluble only to a very small extent. All the acetates smell of acetic acid, on the affusion of sulphuric acid. Those formed with oxides of the metals proper, yield their acid on the application of heat, with a partial decomposition. When the base is a fixed alkali or alkaline earth, they are resolved into carbonates and acetone (3098).

5208. When in diluted aqueous solution, especially when the base is in excess, any alkaline acetate undergoes a

partial resolution into a carbonate.

5209. Of course, any of the acetates may be formed by the saturation of the acid with the proper base. In some cases, they may be obtained advantageously by double de-

• Dr. Ure alleges, that by means of twenty to thirty pounds of platinum powder, which does not waste, we may transform, daily, three hundred pounds of bad spirits

**into the finest** vinegar.

The process has been conducted in a wooden box having a capacity of twelve cu-

the foot

For this purpose, platinum black may be made, by fusing platina ore with twice weight of zinc; pulverizing the resulting alloy, and subjecting it successively to diluted calphuric and diluted nitric acid, the latter with heat. The zinc being disprived or oxidized, the residual powder, after washing with a solution of potash and water, is fit'for the purpose in question.

composition, as illustrated in the case of sulphate of interest and acetate of lead, which when added together in a state of solution, form sulphate of lead and acetate of solution. Formerly, the acetate of potash was known as foliated earth of tartar, acetate of ammonia as spirit of minds rerus.

5210. Of Acetate of Ammonia, or Spirit of Mindows. This salt may be obtained by distilling sal ammoniac villacetate of soda, when, after the escape of some ammoniate the acetate comes over liquefied, and crystallizes in transparent, colourless needles.

5211. This acetate has an acid reaction, is deliquescent and soluble in all proportions in water and alcohol. It the acetates of lead some mention has already been made

(1740.)

5212. Sugar of lead, according to Liebig, contains equal number of atoms of acid and base. Besides the there is sesquibasic acetate, consisting of two atoms of acid with three atoms of base.

5213. Tribasic acetate, consisting of one atom of ad

with three atoms of base.

5214. Sexbasic acetate, holding one atom of acid to atoms of base. Sugar of lead is, of course, the neutracetate.

#### Of Lactic Acid.

5215. This acid is that which exists in sour milk, whence its name from lac, the latin for milk. It has lately been shown to be generally the product of a peculiar fermentation, called viscous, to which the juices of plants, containing albumen, are spontaneously liable, when yeast is not added, at a temperature between 86° and 104°. This immentation differs from the vinous, in being accompanied by the evolution of inflammable gases, as well as carbon acid, and in not being productive of alcohol, but of lactic acid and manna sugar, or mannite (4074). It is obtained from sour milk by saturation with soda, and decomposite the resulting lactate by sulphuric acid. By a previous and dition of lactin, in the ratio of eight ounces to eight put of the milk, the quantity of acid produced may be advantageously increased.

5216. Lactic acid is monobasic, and as it exists in the anhydrous salt which it forms with zinc, consists of C'H'0.

and oxygen which it contains exist in the proportion for forming water, it might be represented as a hydrate of carbon; a composition which usually belongs to bodies, which, as it respects basic and acid properties, act indifferently. When in its most concentrated form, it appears as a sour syrup, incapable of crystallization. On being heated to 482°, it is decomposed, yielding, among other products, a large amount of a crystallized acid sublimate. As this consists of C<sup>6</sup> H<sup>4</sup> O<sup>4</sup>, it was for some time treated as anhydrous lactic acid; but as the anhydrous lactate of zinc is said to contain H<sup>5</sup> O<sup>5</sup>, this sublimate must be regarded as a distinct acid. By boiling in water, the new acid combines with an atom of oxygen and an atom of hydrogen, and is consequently reconverted into lactic acid.

## Of Citric and Malic Acid.

5217. The name of citric acid indicates its origin.\* It exists in the lime and lemon, in union with mucilage and malic acid. Its combination with mucilage is so intimate as to render it impossible to separate the acid without first uniting it with some other matter. Alcohol combines with the acid, and precipitates the mucilage. Yet, the alcoholic solution, thus obtained, does not yield crystals, even after evaporation, re-solution in water, and evaporating the water.

5218. The most efficient mode of obtaining this acid pure, is to saturate the juice of lemons with chalk or whiting, and afterwards to decompose the citrate of lime thus formed, by sulphuric acid, duly diluted. The citric acid may be obtained in crystals, from the supernatant li-

quid, by evaporation.

by acid when concentrated, but agreeably sour when dilute. 5220. It is a tribasic acid; its formula, when dried at 212°, being represented by C<sup>12</sup> H<sup>5</sup> O<sup>11</sup> + 3HO. The atoms of water are essential to the composition of the acid in its free state, and cannot be removed unless by substitution of an equivalent number of atoms of some other base.

5221. Malic acid derives its name from the apple, as

<sup>•</sup> From the fruit of the genus citrus, including the orange, citron, lemon, lime, and sheddock.

in this 'fruit it predominates, as well as in gooseberries, currants, and other similar fruits. It may be had purely saturating lime with apple juice, and decomposing the late of lime by sulphuric acid.

5222. Professor Wm. Rogers, of the University of Virginia, has ascertained that this acid abounds in different species of sumach, in the state of bimalate of lime. Malic acid is bibasic, its formula being C<sup>5</sup> H<sup>4</sup> O<sup>5</sup> + 2HO.

a law, to which a great many of the vegetable acids are subjected. At a perature a little above that at which they melt, they severally yield new acid. That yielded by citric acid, is identical with the acid found in the aconim napellus, and also the various species of equisitum. Hence, it has received the name of aconitic or equisitic acid. Whether obtained from citric acid by heat, or from either of its other sources, it exists in the form of which acid is changed, under similar circumstances, is also found in nature in the Iceland moss, and in the fumaria officinalis. Hence it has been called in maric acid, although Pelouze, who first obtained it from malic acid by heat, called it paramalic acid. Both of these acids differ from the citric and malic acid, from which they are produced, only in having lost the element of two atoms of water.

5224. When either of the acids thus obtained, by heating citric or mix acid, is exposed to a higher temperature, a further change takes place, will volatile acids are formed, fumaric acid yielding malic, and aconitic producing itaconic acid. The former would seem to be formed by a mere transposition of the elements of water present, which appear as two atoms of water of crystallization, instead of entering as before as two basic atoms into the integral composition of the acid. A farther application of heat converts itaconic into citraconic acid; while malleic acid, if kept in a state of fusion for a length of time, reverts to the condition of fumaric acid.

keeping them at the temperatures necessary for the formation of the socioompounds which they respectively produce, the result will be a mixture is the one case of fumaric acid and malic acid, in the other, of aconic, is

conic and citraconic acids.

## Of Tartaric Acid, and Paratartaric or Racemic Acid.

5226. Tartaric acid is found in many vegetables. It is named from tartar, an appellation given to a deposition from wine, which contains this acid united with potash and water. This tartrate, when freed from impurities, is known officinally under the name of cream of tartar. When to twenty-four parts of this salt, thirteen of carbonate of soda are added, sal Rochelle, a tartrate of potash and soda, is produced; and in like manner, tartar emetic, by replacing the basic water by the sesquioxide of antimony. Another pharmaceutical compound, called tartarized iron, is pre-

ced by replacing the water of cream of tartar by iron, ich is taken up in the state of protoxide, but becomes,

exposure, more or less sesquioxidized.

5227. Tartaric acid is procured from cream of tartar in secrystals, by adding chalk until effervescence ceases, decomposing the precipitate by diluted sulphuric acid. he neutral tartrate of potash left, may be decomposed by sicklime or chloride of calcium, and the resulting tartrate lime will yield the acid in the same way as the analous tartrate, obtained in the first instance by the addition chalk.

5228. Tartaric acid is extremely sour, and reddens litus. It is bibasic, its formula being C<sup>8</sup> H<sup>4</sup> O<sup>10</sup> + 2HO.
I consequence of this bibasic character, the salts which it
rms with one atom of a fixed base are sour, have an acid
action, and require the presence of an atom of basic
ater. Thus the salt heretofore described as the bitarate of potash, must now be considered as the tartrate of
stash and water, since it consists of one atom of tartaric
id, one atom of potash, and an atom of basic water.

5229. Of Paratartaric or Racemic Acid.—A manufacturer of Thann, in Germany, in preparing tartaric acid om cream of tartar, which had been deposited from the ine of that country, discovered an acid differing from lat which it was his object to procure, and which he supposed to be the oxalic. Gay-Lussac subsequently proved, lat while possessed of peculiar qualities, its equivalent as the same as that of tartaric acid. By Berzelius it as afterwards shown to be isomeric with this last menoned acid, and he has consequently named it paratartic acid. The appellation of racemic, has also been apied to it. Paratartaric acid crystallizes in a different rm from tartaric acid proper. It is likewise less solu-

5230. The action of heat on tartaric acid is strikingly peculiar. At a meaning merely sufficient to produce fusion, two atoms of the acid give fone of the four atoms of the basic water combined with them, losing at same time one fourth of their saturating power, and causing the acid to come sesquibasic, so that two atoms of it saturate only three of base. he name of tartralic has been applied to the acid in this state.

5231. A still further application of heat removes another half atom of ster and produces tartrelic acid, which is monobasic, saturating only one of base, and requiring in the free state the presence of but one atom water. A still higher temperature removes all basic water, and leaves a rous white mass, insoluble in water, and hence no longer sour or capable

of reddening litmus. The composition of this body is C<sup>o</sup> H<sup>o</sup> O<sup>10</sup>. Case quently, it is identical with that of tartaric acid freed from its basic war, as it exists for instance in the bibasic tartrate of lead. If left long in catact with the water, this insoluble compound gradually takes up two stors of the oxide of hydrogen, and becomes the ordinary soluble bibasic tarance It has been considered, that the absence of sourness, in this only stance of an isolated anhydrous organic acid, is favourable to the idea that oxacids are hydrurets of compound radicals owing their acid reaction to hydrogen; but it should be recollected, that the absence of this action is an invariable consequence of insolubility. No insoluble hydruret of which there are instances among the oils or etherial compounds is sour. It is it that portion of water which enters the tartaric acid as a base, and cabe hydrogen of which the hypothesis relies, which confers either sources er Independently of the capacity for acid reaction with vegetable colours. moisture, the gaseous hydracids, erroneously so called, have, I believe, D such properties.

5232. Of Liquid and Solid Pyrotartaric Acid.—By destructive distinction, tartaric acid yields two acids, to which the preceding appellation have been given. Liquid pyrotartaric acid forms a monobasic ether, and various salts. Its formula is C<sup>6</sup> H<sup>2</sup> O<sup>5</sup>. Solid pyrotartaric acid is granted in small proportion, during the destructive distillation of tartaric acid but is yielded more copiously by subjecting cream of tartar to that process.

Graham, 948.

#### Of Guiacine, or Guiacinic Acid.

5233. In the Journale de Pharmacie, for 1842, p. 386, notice is given by J. Pelletjer, of the results of an investigation, which, though it had not been completed, enabled him to allege that the peculiar principle of gum guinces, which he calls guayacine, in English guiacine, may be isolated by either of two processes. According to one, an alcoholic solution of accuse of lead is to be added in successive portions to a tincture of the resin. rejecting the latter portions of the precipitates formed. The compound thus procured, is to be well washed with water first, and afterwards with alcohol. Then being suspended in water, is to be exposed to sulphydric acid, by which the lead is precipitated as a sulphide. The guiacine is then taken up by alcohol.

5234. According to the other process, hydrate of lime is added to the tincture, by which means a compound, of the guiacine and lime, is obtained.

From this the guiacine may be easily extricated.

5235. Guiacine has, in a high degree, the property of becoming blue is absorbing oxygen, and, after being thus coloured, may be restored to previous state by substances greedy of oxygen, such as sulphydric or phurous acid, protoxide of iron, or protochloride of tin. Re-exposure to the air restores the blue colour.

5236. Moist chlorine, or an aqueous solution of this gas, turns guiscine blue; but an excess renders it green, and yellow, successively. From the last mentioned state it cannot be restored, having undergone a chemical change.

5237. Notwithstanding the property of combining with bases. Mr. Pelletier hesitated to designate it as an acid, but in this, as it was found to combine with bases, I consider him as misjudging. The analogy between the resin and indigo, as respects changes of colour, must strike every one sequainted with the facts.

5238. I presume in English the principle which he has isolated, will be led guaicine; or if it be an acid, as from the account given, it evidently ght to be considered, the name will be guaicinic acid.

## Of Tannic Acid.

5239. From its formula, C<sup>18</sup> H<sup>5</sup> O<sup>9</sup> + 3HO, it may be en that the tannic acid is tribasic.

5240. The art of converting the hides or skins of anials into leather, by soaking them in infusions of the bark oak and other trees, had long been practised. Subsemently it was ascertained that this change arose from a cheical combination ensuing between the gelatin of the skin rhide, and a vegetable principle called tannin, from its efciency in the process of tanning abovementioned. Berdius first treated of tannin as an acid. This view being lopted, the principle is now universally designated as nnic acid. It is peculiarly abundant in oak galls, giving an infusion of them the property of causing, with iron, ink colour, whence its use as an ingredient of common riting ink.

5241. Tannic acid is likewise found in a great number vegetables, generally in their bark or roots, but not unequently in their leaves and seeds, and even in their flows and fruits, before they have reached maturity. It is, fact, the most frequent cause of astringency in vegetable

5242. It may be procured, according to Mr. Pelouze, in state of purity, by introducing powdered galls into a vesl, with a body and pipe resembling that of a funnel, but intracted above into a neck like that of a bottle. The pe of this vessel should be furnished with a cock, and

ust be made to descend into a tincture bottle through e mouth. The galls are then to be covered with suluric ether, of the officinal strength, and the mouth of the seel being corked, they are to be left in contact with the her for several hours. The liquid being then allowed to secend into the bottle, will be found to separate into two ortions, of which the heaviest is a solution of tannic acid. rom this solution the acid may be obtained in the solid rm by washing with ether, and evaporation, in vacuo, rer sulphuric acid. Thus obtained, it is inodorous, as-

ingent, yellowish white, and somewhat crystalline.
5243. The oxides of the following metals form insoluble

tannates, and hence yield precipitates with tannic acid, an infusion of galls. The colours of these precipitates as follows:—

The precipitate formed with lead or antimony, white.

With tantalum or bismuth, orange.

With titanium, blood red.

With platinum, green.

With chrome, molybdenum, uranium, and gold, brown With osmium and sesquioxide of iron, deep purple, or ink colour.

5244. On account of the insolubility of the tanks antimony, an infusion of galls, or of oak bark, is at dote for tartar emetic and other antimonial preparating

5245. Tannic acid has also been found a test for precipitant of, the organic alkalies, and must be more

less an antidote for their poisonous influence.

5246. The aqueous solution of tannic acid reddent mus. It does not affect solutions of the protoxide of and the intense colour produced as abovementioned, the sesquioxide, may be removed by reagents, which duce the iron to the state of protoxide, as already trated (1817).

5247. Ink is best made with the green sulphate of because, so long as the iron is not sesquioxidized, reing in solution, it can penetrate the paper better; at a soon peroxidizes, and consequently blackens, by exposure

to the atmospheric oxygen. (Urc.)

5248. By a piece of raw hide, pure tannic acid may, a few hours, be taken up from a solution so completely that if no gallic acid be present, the liquid will not be fected by a solution of sesquioxide of iron.

5249. According to Graham, tannic acid precipitated solution of starch and albumen, and is capable of

bining with animal fibrin.

tannic acid in many of its properties, and called, generally, artificial tannin, is formed during the action of its tric or sulphuric acid on a great variety of vegetable stances. One variety of this tannin is formed by the restation of nitric acid with charcoal.

## Of Gallic Acid.

1. Formula of the dry acid, C<sup>7</sup> HO<sup>3</sup> 2HO. When lized, one additional atom of water is present.

2. This acid and tannic acid appear to be almost

more or less associated; so that they are generally resent, where either is found. This is now explained fact, that tannic acid is liable to be converted into

acid spontaneously.

- 3. Agreeably to one of the processes recommended curing the last mentioned acid, nut galls, made into with water, are to be exposed to the air for seve-eks at the temperature of 80° nearly, water being d so as to compensate for evaporation. The remass is to be subjected to boiling water, and the n thus obtained being filtered, the gallic acid sepanthe crystalline form. It is rendered quite pure by tion, digestion with animal charcoal, and re-crystal-
- l. If the precipitate, obtained by adding sulphuric a concentrated extract of galls, be washed with a quantity of water, and then dissolved by gradually it to a boiling solution of one part of sulphuric acid of water, gallic acid is generated, and, by refrige-separates from the liquid in crystals. The impure us isolated, may be purified partially by re-solution ystallization; or more thoroughly by adding to a n of it acetate of lead, and decomposing the result-oluble gallate of the protoxide of lead, by sulphydric lead, which separates this metal, and much colouring, simultaneously; the acid remaining dissolved. n. 941.
- i. Again, if tannic acid be subjected, for a few mito a solution of caustic potash, on the addition of ric acid in excess, crystals of gallic acid will be coformed on the cooling of the liquid. Kane, 1010.

  i. Gallic acid crystallizes from a hot solution in thin eedles, which, for solution, require 100 parts of cold although, when boiling, three parts are sufficientary soluble in alcohol, and sparingly soluble in ether. gh it is productive of the same changes in solutions

of sesquioxide of iron as tan , it di iers from it not causing any precipitate in solutions of gelatine.

5257. It would appear doubtful whether this acid ready formed in nature, or whether it be not always and duct of the oxidation, or partial decomposition of acid. It has been stated, that the exposure of the to the air, or boiling it with an excess of alkali, the presence of the atmosphere, produces this character and that it may also be effected by means of supposed.

5258. On the one hand it has been observed, that atoms of tannic acid contain the elements of six ato gallic acid, and one of grape sugar; and on the other, the absorption of eight atoms of oxygen would conve atom of tannic acid into four atoms of carbonic acid two of crystallized gallic acid. As, according to Br not, alcohol and carbonic acid have been evolved from galls during their fermentation, it seems possible that nic acid may be produced, according to circumst either by fermentation, or by the oxidation of the ples present in nut-galls. Indeed, tannic acid itself appear, from the nature of the sources from which obtained, to be, in many instances, the result of a gr decay of other principles in plants; and when gallic either in its free state, or as it exists in the gallates, is conposed to the air, it undergoes a still further change into carbonic acid, and a brown vegetable substance. it may be conjectured, that both of the acids in question are the products of different stages of one continued trasformation.

bonic acid, and a new acid which sublimes in brilliant white plates. This acid has received the name of pyrogallic, and is soluble in water, and ether. If, on the contrary, the heat be raised above 450°, an install black mass remains in the retort, to which, from its combining with the lies, and its colour, the name of melangallic acid has been given. The results are only worthy of notice as forming part of a series of transformations which most of the organic acids undergo through the application heat.

5260. An acid, called the elagic, is frequently produced during that a posure of galls to the air, which gives rise to the formation of gallic and the several species of vegetable products in which acids, resembles the gallic and tannic acids, though not identical with them, have been covered. Thus, in the bark of the various species of cinchona, combined with quinia or cinchona, are found two acids, the cinchonic and cinchonal cinc

nic, whose physical properties stand in very nearly the same relation to h other as that borne by gallic and tannic acid; and in catechu, an ext obtained from the mimosa catechu, there have been discovered two ds, the catechuic and the catechutannic, of which nearly the same statent may be made. It does not appear, however, that in either case one them has the property of being converted into the other, as is the case h tannic and gallic acids. Berzelius, however, is of the opinion, that all forms of tannic acid found in plants are identical in composition, but dified by association with other matter.

## Of Meconic Acid.

5261. Formula, C<sup>14</sup>HO<sup>11</sup>+3HO; when crystallized +6HO. econic acid is tribasic.

5262. When a solution of acetate of lead is added to an usion of opium, a precipitate is obtained, consisting of conate of lead. From this the lead may be precipitated a sulphide by means of sulphydric acid, and a solun of the liberated meconic acid obtained by filtration is acid exists in opium, combined with morphia and leia.

5263. With solutions of the sesquioxide of iron, meconic d produces an intense red colour; with protoxide of lead insoluble precipitate. It is to this affinity, for oxidized d, that we owe the process, above described, for proring this acid.

5264. Meconic acid produces a taste, at first sour, and sequently bitter, and reddens litmus paper. Being a pasic acid, it forms three classes of salts, in which the ter present may be replaced, partially or entirely, by e, two, or three atoms of base. Like other organic ds which have been described, meconic acid is converted heat into another acid, the komenic, carbonic acid being lived; and as this komenic acid cannot be volatilized, it at a higher temperature, converted into pyromeconic d, which may be sublimed without further change. ch of these transformations is accompanied by the loss an atom of basic water, and a diminished capacity of urating bases.

## Of a Method of detecting the Presence of Opium.

5265. The property which meconic acid has of precipiing with lead, and of producing a red colour with iron, y enable us to detect opium, when present in a very all quantity in solution.

5266. If ten drops of t tincture ( called laudanum, be min; d with half a gallon of wa on adding a few drops of bate of lead, there will precipitation which, at t e f a few hours, will be ceptible in flocks. To de tof these flocks may celerated by detachi t gently from the sides recipient with a g . I. The vessel should be c so as to concent: t during their descent. they are collected at t pottom of the vessel, about drops of the red sulphate of iron, and an equivalent porti of sulphuric acid should be introduced among them means of a small glass tube. The presence of the conic acid will be rendered evident by the redness which ensues.

5267. When a red colour is produced by the manhere described, it is probable that opium is present; as conic acid is found only in that drug, and having no acqualities, is not used separately from it in any pharmatical preparation.

5268. It may be proper to mention, that sulphocyands dric acid produces, with the sesquioxide of iron, a color

resembling that produced by meconic acid.

## Of the Acids formed from Sugar.

5269. Cane sugar may be made to combine, as sugar, with the alkaliance earths, and with some of the metallic oxides, though not with the alkaliance. In these compounds the sugar exists unchanged, but united to the base by

an affinity so feeble, that it may be displaced by carbonic acid.

5270. Nevertheless, if sugar be kept a long time dissolved in an alkalian solution, it undergoes a transformation into a real acid, the glucic which has a sour taste when free, and combines with bases to form substantially as in lactic acid, the oxygen and hydrogen are present in the proportion for forming water; and the only change which sugar experience by conversion into glucic acid, is the loss of several atoms of water. The formula for glucic acid would appear to be C<sup>12</sup> H<sup>8</sup> O<sup>8</sup>.

5271. If heat be applied to a solution of sugar with an alkalise melassic acid is produced either from the sugar directly, or from the paid acid. It is said to consist of C<sup>24</sup> H<sup>13</sup> O<sup>10</sup>, so that in forming it, sugar paids

not only with water, but also with oxygen.

5272. By the reaction of diluted nitric acid with sugar, a crystalliable acid of a strong sour taste is produced. It was at first supposed to be said, but was afterwards distinguished by the name of oxalhydric. It is now called saccharic acid. Its formula is C<sup>18</sup> H<sup>5</sup> O<sup>14</sup> 5HO. The first atoms of water are essential to the composition of the acid in what is called the free state. When it is united to other bases, the water is replaced, wholly or in part, by a corresponding number of atoms of base. The same hydrous salt which it forms with lead, consists of C<sup>18</sup> H<sup>5</sup> O<sup>11</sup> + 5PIO:

d by its union with the oxide of that metal, it forms three other salts, in hich we find C<sup>13</sup> H<sup>5</sup> O<sup>11</sup> combined, respectively, with 3PbO + 2HO, 2PbO 3HO, and PbO + 4HO. These facts respecting the composition of the ccharates are instructive, as furnishing support to the theory of polysic acids; since, if we do not have recourse to that theory, we must suppose the existence of a distinct acid in each of the salts above mentioned, and at one of them has the property of combining with five atoms of base, and at with any smaller quantity.

5273. When lactin (sugar of milk, 4070) is subjected to the action of luted nitric acid, mucic acid is produced. It may also be obtained by betituting gum, or mannite, for the sugar of milk. It exists as a crystal
ne powder of difficult solubility, and a feebly acid taste. Its formula is

49  $H^{\circ}$  O<sup>14</sup> + 2HO.

## Of Formic Acid.

5274. It is inferred, that between formic acid, formyl 4019), and methyl (4016), the same relation exists as beween acetic acid, acetyl, and ethyl; and also that the part erformed by alcohol, the hydrated oxide of ethyl, in the ne case, is performed by pyroxylic spirit, the hydrated xide of methyl in the other. Either the methylic, or ethycalcohol, by losing two atoms of hydrogen, and acquiring wo of oxygen, are converted, the one into acetic, the other to formic acid. Moreover, the same catalytic agent, planum sponge, or black, may in either case be competent induce the requisite reaction with atmospheric oxygen. The features which are wanting to complete the resemlance, are congeners severally of aldehyde, C4 H3O + HO, acetous acid, C4 H3O2 + HO. To correspond with ness compounds, no hydrated oxide of formyl, nor formous cid, are known.

5275. To render this statement more intelligible, the sllowing formulæ are subjoined. Methyl, C<sup>2</sup> H<sup>3</sup>; formyl, H; anhydrous formic acid, C<sup>2</sup> HO<sup>3</sup>. To form the hyrated acid, one atom of water, HO, must be added.

5276. Formic acid was originally obtained from ants.

: appears to exist in them naturally.

5277. This acid may be obtained by adding to one art of sugar in an alembic, three parts of well pulverized eroxide of manganese, and three parts of sulphuric acid iluted with its weight of water. The acid should be dded in three successive portions. At first, the effervesence is so great as to require the vessel to have fifteen mes the capacity which would be necessary to contain the material when quiescent. The formic acid associated

with formic ether, is brought over by d | ion. It my be saturated with chalk or an alkali, and the results formiate decomposed and isolated by distillation with parts by weight of sulphuric acid, diluted with forward.

5278. According to the late Professor Emmet, the sence of peroxide of manganese in this process is used sary. Agreeably to his observations, the conversion many vegetable substances into formic acid, among of maize, may be effected by any of those agents who would effect the evolution of ether from alcohol.

5279. From the investigations of Dobereiner, it a pears that formic acid is an excellent reagent for separing the noble metals from solutions in which they are termingled with other metals proper. If a solution taining one or more noble metals, be elevated nearly test temperature of ebullition, on adding an alkaline formit the noble metals will be immediately and entirely pretitated in a very minute state of division. At the same the by ascertaining the weight of the gas simultaneous evolved, that of the metal thrown down may be determined.

5280. From its solution in water, the bichloride of macury is converted into calomel with so much facility, in a state of division so perfect, by formic acid or formitted of soda, that their employment in the preparation of that

protochloride was suggested by Dobereiner.

5281. If the same quantity of sulphuric acid and manganese be mingled with six parts of alcohol, the process being, in other respects, the same as that for formic acid above described, formic ether becomes the predominant product. It is freed from formic acid by magnesia, from alcohol by a small quantity of water, and from water by chloride of calcium. By a more extensive contact with water, formic ether is decomposed, and alcohol and formic acid are generated.

5282. Formic acid has a pungent taste, and a peculiar sharp odour. It is more energetic in its affinities than acetic acid. The formiates, like the acetates, are generally

rally very soluble.

#### Of Valerianic Acid, C10 H9 O3 + HO.

5283. This acid was described in the last edition of this Compendium, a product yielded by the root of valerian, (valeriana officinalis,) when bjected to distillation with water. Since that time, it has been found to producible, artificially, from a totally different source. It has been disvered by Cahours, that if oil of potato spirit (hydrated oxide of amyl, 123), be allowed to fall in successive drops no faster than it can be imbed upon platinum black, previously heated, an acid vapour arises from exidation of the elements of the oils, which has all the properties of valerianic acid, obtained from the root of valerian as above mentioned. 5284. During this process, two atoms of hydrogen are replaced by two oms of oxygen, so that it is quite analogous to the play of affinities by hich the acetic and formic acids are generated; the former from alcohol, latter from pyroxylic spirit.

5285. Valerianic acid is also generated in potato spirit, by the sponta-

sous absorption of atmospheric oxygen by exposure to the air.

5286. Valerianic acid is a colourless liquid, having an oleaginous constency, a sharp, acid taste, and a persistent odour, which recalls that of root of valerian. In the state of protohydrate, according to Graham, produces a white spot upon the tongue when applied to it. The density this acid is nearly 937 at 62°. It boils without alteration at 347°, and mains liquid at 5°. When heated in a platinum spoon it takes fire really, burning with a white flame and much smoke, leaving little residue. is soluble in eighty times its weight of cold water, and in all proportions alcohol. It is capable of taking up 20 per cent. of water without losing oily consistence.

5287. From the formula of this acid it is supposed, that it may consist a compound radical analogous to acetyl, for which the name valeryl is

ggested; formula, C10 H9.

#### Of Caffeic Acid and Caffee Tannic Acid.

5288. According to Kane, the coloured precipitate produced in a decocnon of raw coffee, by subacetate of lead, comprises two substances, which may be extracted by impregnation with sulphydric acid while suspended in ater, subsequent evaporation of the filtered liquid to the consistence of rup, and digestion in strong alcohol. A peculiar kind of tannic acid dislives, called caffee tannic. A white powder subsides, which, when heated, rolves the peculiar smell of roasted coffee. Its solution in water reddens mus. It is called caffeic acid. It is not known whether the tannic acid tea and coffee are the same.

## Of Acids modified by an Union with Organic Matter.

5289. Two sets of acids may claim this description. If these, in one set the organic matter to which the hange in them is due, is an oxidized compound organic idical, acting as a base, capable, under favourable circumtances, of being transferred to other acids. In the other it, the matter producing the change does not contain a

compound radical, capable by oxidation of acting and base, and transferrable to other acids.

## Of Acids modified by Union with an Oxidized Computer Radical.

5290. Acids of this set. ien formed of a mondance acid, require for existence atoms of acid and two stems of base. One of the acid is must be in union with the oxidized radical, the other union with an atom of basic water, or some other oxide g as a base. Hence, 25 already suggested in the ca sulphovinic acid (3000) as double salts of an on auch compounds may be vie dized radical and oxide of hydrogen, so that, agreeably 10 the language of Graham, sulp winic acid is a sulphate d other and water. But this d s not explain the fact, that and oxidized radical canad a *neutral* compound of the be made. Hence. v, presented by the same author, seems to isfactory, agreeably to which re one bibasic acid, of comme the two atoms of a 1 it has been formed. This reisomeric with that of tionale seems to derive strength from the fact, that 🚥 atom of tartaric acid in tartrovinic acid, performs the patt of two of sulphuric acid in sulphovinic acid, agreeably w the usual idea.

5291. In the other set of modified acids, the organic matter does not appear to be in a basic state, not being an oxide of a compound radical, nor capable of separation without decomposition.

5292. In three of the acids belonging to the first st (sulphovinic, phosphovinic, arseniovinic acid), ethyl, being the principal radical, is united to an inorganic acid.

methylic acid, in which the oxide of a compound radical plays the same part in combination with a double atom of sulphuric acid, that the oxide of ethyl plays in the three acids above mentioned. Also in tartrovinic, oxalovinic and camphovinic acid, one atom of tartaric, oxalic, or camphoric acid, performs the office of a double atom of sulphuric, arsenic, or phosphoric acid, in the analogous compounds arising from their association with the same oxidized radical. Other acids exist, having a similar composition of the same oxidized radical.

tution to those last mentioned, and that there will be iny more produced hereafter, there is much reason to ppose.

5294. In the second set, there are several which are asbed to an union of hyposulphuric acid with carbon, hy-

ogen, and water; as, for instance,

\*\*phosulpho-naphthalic sposulpho-benzoic acid consisting of S<sup>2</sup> O<sup>5</sup> hyposulphu-  $+ C^4 H^4 O$  ric acid  $+ C^{14} H^4 O^3$ 

5295. Other acids consist of the elements of some defite organic compound, such as sugar or indigo, so united an acid as to form, with bases, crystalline compounds; iich, besides the peculiarity of their crystalline form, ve a solubility altogether wanting in the salts generated, r se, by the acid with which they are formed. This deription is intended for sulpho-saccharic acid, and hypolpho-indigotic acid; one created by the union of indigo th hyposulphuric acid, the other by the union of sugar th sulphuric acid.

5296. Analogous to the former of these acids, a new id has been created, by the reaction between sulphuric id and acetic acid, of which the formula, represented

C' H' O' + S' O', makes it a compound of hyposulurous acid; but Berzelius suggests that the same ultite elements, in a different order, would give C' H' O' 2SO';\* and that the formula thus made out, being dided by two, would give C' H'O + SO'. This would make a sulphated oxide of elayl, of olefiant gas in other words 095).†

Of Sulphorinic Acid, or the Sulphate of Ether, and Water.

5297. Of the acids above described, of the first class, I shall here treat sulphovinic acid only. While the limits prescribed to a text-book do not ow me to do more, the importance of this acid, arising from the part ich it performs in the production of ethers, and the expediency of select; it as an exemplification of the set to which it belongs, renders it prothat I should add something to the notice already taken of it under the ad of ethyl (3069).

5298. Sulphovinic acid is produced by heating, to the boiling point of the ulting mixture, or about 280°, equal weights of concentrated sulphuric d and alcohol of from 830° to 850°; or by saturating sulphuric acid with vapour of ether, and adding water after some hours have elapsed. In

Report on Chemistry for 1841.

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It also contains the elements of a hydrated bisulphate of the oxide of acetyl, H<sup>2</sup>O + 2SO<sup>2</sup> + HO.

the case, the resulting liquid is to be esturated wind of had, the lime, or baryta. From the resulting sulphovimes, the sulphovime the may be liberated by adding enough acid to saturate the inorganic base, at in the case of that formed with lead, by precipitating this metal by subjections.

dric acid (899).

5290. It has been mentioned that sulphovinic acid is equivalent to a ble sulphate of ether and water; but that the two storm of sulphane of act like one atom of a peculiar bibasic acid, isomeric with sulphane and since it cannot be obtained as a neutral compound, consisting of one was of bride of ethyl and one of anhydrous sulpharic acid. The heavy of wine, heretofore considered as a neutral hydrated sulphate of ether and ethers, of atheroic agreeably to a new name employed by Liebig.

5300. Any stronger base presented to sulphovinic acid takes the placed the water, as in other cases where bibasic acids are in union with as also of water and an atom of another base; but the acid in question differs from where bibasic acids in this, that the oxide of compound radical is essential the chautance of its bibasic property. As soon as the oxide of ethyl was placed, the monobasic character of the inorganic acid is resemmed.

5301. The suits formed with sulphovi ic acid by the replacement of the basic water, have been known as sulphovinates; and so long as the suits of the acid is debuteable, it will be preferable to adhere to this name.

330%. One of the most remarkable traits of the sulphovintres, is that a lability which prevents the detection of sulphuric acid, even by solution a

beryta.

by their distillation with alcohol and sulphuric acid, it is probable that the formation of sulphovinic acid is a preliminary effect (3006). But when the mixture is subjected to heat, the organic acid and oxide of ethyl, being competent to form a volatile compound, go off in union, while the minary to tween the water and sulphuric acid co-operates to expel them. It is, therefore, a case of double elective affinity, aided by the vaporizing influence of caloric.

8804. It was upon this view of the subject, that Messrs. Boyé and Homewere enabled to foresee the production of the wonderfully explosive percharic ether, by distilling perchlorate of baryta with the sulphovimus of the same base.

#### Of Succinic Acid.

sublimed in crystals, much contaminated by the essential oil of amber. It digestion in nitric acid, evaporation to dryness, washing in cold water, sequent solution in boiling water, and finally by crystallization, the soul obtained pure. When combined with any of the alkalies, it is useful in parating the sesquioxide of iron from the oxide of manganese.

5306. Succinic acid has a sour taste, and reddens litimus paper. To formula of this acid, in the hydrated state, is C<sup>4</sup> H<sup>2</sup> O<sup>3</sup> + HO, being from the formula monobasic. The formula of the sublimed acid is C<sup>4</sup> H<sup>2</sup> O<sup>3</sup>.

#### Of Benzoic Acid, C14 H5 O4 + HO.

5307. From the formula it must be evident that this is a monobasic soil containing one atom of oxygen more than the compound radical, beneals, C. H. O. (3063): also, that replacing an atom of oxygen by one of by:

ogen, must convert it into the hydruret of that radical, C<sup>14</sup> H<sup>5</sup> O<sup>2</sup> + H

120). The symbol of this acid is Bz.

5308. Benzoic acid exists ready formed in the resinous product of vegeion improperly called gum benzoin, from which it may be extricated by
following process:—Spread a pound of the benzoin over the bottom of a
st iron pot, of eight or nine inches in diameter and of about two inches in
pth, the mouth of the pot being covered by filtering paper secured to the
m by paste. Thus covered, the pot is to have a canopy of coarse packpaper fitted to it, like a cap, and secured by a bandage of wire. The
t, thus charged and prepared, is to be subjected to a sand bath for three
four hours. Under these circumstances, the cavity, included between the
ver and cap, becomes studded with crystals of benzoic acid, quite free
m the black empyreumatic oil with which they are liable to be soiled,
ten sublimed without the interposition of the paper. The acid thus extrited, amounts usually to about four per cent. of the gum employed.

5309. Benzoic acid may likewise be obtained by boiling, in four parts of iter, equal parts of gum benzoin and hydrate of lime, until the liquid is luced to 4th of the original volume. It is in the next place to be filtered, d an addition made of chlorohydric acid. After a second filtration, the

id separates in crystals.

5310. It is presumed that benzoic acid exists in the gum ready formed, d is therefore liable either to be sublimed by heat, or removed by its inity for lime, from which it is expelled by the chlorine of the chlorohy-c acid.\*

flexible laminæ, white, pearly and translucent. When pure it is inodous, though by being heated, it acquires a smell analogous to that of bennalowed. It reddens litmus feebly, melts at 248°, and sublimes at 293°, osphorescing in the dark. It boils at 492.2. Heated in the air it yields very acrid white vapour, which excites coughing. It is highly inflammation, burning with a very smoky flame, leaving no residue. It is soluble 100 parts of cold water, and in 25 at the temperature of ebullition; varizing with its aqueous solvent when this is distilled from it. For solution a part requires two parts either of ether or alcohol.

5312. Peroxide of iron precipitates in the form of an insoluble subbenate, of a reddish white or buff colour, when a soluble benzoate is added a solution of peroxide of iron, previously neutralized by ammonia withtany consequent precipitation. Hence benzoate of ammonia serves to parate the sesquioxide of iron from the oxides of manganese, nickel, or ic; when the solution contains neither alumina, yttria, zirconia, nor glua; of either of which the oxides would be simultaneously precipitated if

ment. Graham, 851.

# Of Hippuric Acid, C13 H8 N2 O5 + HO.

5313. Hippuric acid is found in the urine of herbiverous mammalia. ebig supposes that it is probably derived from food, in which it pre-exists;

M. Jahn has remarked that when the sawdust of guiacum wood (lignum vite) is ated with a solution of carbonate of soda, sulphuric acid added, the liquid and the in which precipitates is washed, dried, and subjected to heat in an appropriate apratus, a small quantity of sublimed benzoic acid is obtained. Berzelius' Report, 11, 106. This justifies an inference made by Guibourt, that the crystalline partime observable in the bark of the wood in question, might be benzoic acid.

since it abounds in the urine of horses fed with f bles, but in the of the same animals fed with hay, or other dried vegetable matter, in p-

placed by benzoic acid.

fresh urine of the horse or cow, at a temperature carefully kept below to of ebullition, adding sufficient chlorohydric acid to render it pesses acid, and subsequently allowing it repose. By these means, impaired coloured hippuric acid separates in crystals. It may be purified to pregnation with chlorine, or by the addition of chlorohydric acid and the

ing salt, until the smell and discoloration are removed.

5315. Hippuric acid reddens litmus, crystallizes in large four-aided parent prisms, susceptible of fusion, without loss of weight, into an el nous liquid which yields crystals on cooling. At temperatures higher its point of fusion, it may be decomposed into benzoic acid and be of ammonia, which may be distilled and condensed in red drops, asso with an oily product having an agreeable odour resembling that of Towards the end of the distillation, cyanhydric acid a over, leaving a porous residue of curbon. Hippuric acid requires 460 j of cold water for its solution, but is very soluble in hot water, and still sp in alcohol. In ether it is but slightly soluble. Concentrated acid dissolves hippuric acid without discoloration; but at a higher rature the solution blackens, evolving sulphurous acid, and a sublis beazoic acid. By nitric acid it is transformed into benzoic acid. In e hydric acid it dissolves without alteration. Peroxide of manganese sulphuric acid, aided by heat, convert it into carbonic acid, ammoni benzoic acid. Boiled in water with the puce oxide of lead, it is transf into the amiduret of benzule, or benzamide and carbonic acid. with four times its weight of slaked lime, this acid is converted into nia and a volatile oil called benzole, with a greyish residuum.

5316. When the urine of the horse is left to itself for a long while, at exposed to a rapid evaporation, only benzoic acid is found therein. It is in

combination with ammonia or soda that this acid exists in urine.

5317. Of the Hippurates.—The combinations of hippuric acid with the oxides of metals proper, excepting iron, being more soluble in water when boiling than when cold, may be obtained in crystals, from an aqueous plution made at the temperature of ebullition, and subsequently cooled. By reaction with the hydrates of potash or lime, the hypurates yield ammonia, and an oily liquid, probably benzole.

5318. Of Formobenzulic Acid, Bz H<sup>6</sup> 2HO<sup>8</sup> + HO.—This acid consists of formic acid and the hydruret of benzule, being created during the reaction of chlorohydric acid with the distilled water of bitter almost, comprising, of course, the hydruret and cyanhydric acid (3055). The cyanhydric acid is decomposed into ammonia and formic acid. With the

acid, while nascent, the hydruret combines.

5319. Formobenzulic acid, thus obtained, is in the state of pulverules white crystals, fusible into an oily liquid at the expense of the water of crystalization, and capable, when aided by heat, of decomposing the accuse, carbonates, and benzoates. Its aqueous solution, when submitted to cherine, to nitric acid, or to peroxide of manganese with diluted sulphuric acid, is resolved into carbonic acid, and the hydruret which forms its characteristic ingredient. It has the same capacity of saturation as formic acid. Of course it is a monobasic acid.

#### OF THE ACID OILS OF SPIREA ULMARIA AND GAULTHERIA.

5320. As the results obtained by Mr. Procter, Jr., respecting the analogy r identity of the oils of gaultheria and spirea ulmaria, must create a desire be acquainted with both, I have abstracted, with some changes in the tyle, from Gregory's translation of Liebig, so much as relates to saliculous cid, and have subjoined some quotations from Procter's memoir. thought it necessary to alter the names employed by Gregory. Yet alicyl might be considered as a compound halogen body, combining with ydrogen, like cyanogen in cyanhydric acid, in which case, consistently, s name would be salicohydric acid. Of course this acid, in combining rith oxybases, would have to be considered as generating salicides of their adicals respectively. Were this mode of contemplating the subject admisible, the atom of hydrogen which forms the radical in salicohydric acid, nust be supposed to be converted into water by uniting with an equivalent of oxygen from the oxybase of any radical with which it may combine. nust, however, be evident, that the adoption of these innovations in nomenlature would be attended by great practical inconvenience from the consequent multiplication of discordant names. And were it otherwise, I should iot deem it judicious to make the suggested changes, because our knowedge of such compound radicals as salicyl, and of the state in which they xist in their alleged combinations, is altogether hypothetical and insuscepible of any conclusive proof. Salicylous acid may be viewed as a hydrated racid,  $C^{14}$  H<sup>5</sup> O<sup>3</sup> + HO (3063, 5343).

# Of Salicylous or Saliculous Acid, also called the Hydruret of Salicyl, but more properly considered as Salicohydric Acid, C14 H5 O4 + H.

5321. This acid was first discovered by Pagenstecher in the volatile oil spirae ulmaria; by Piria as a product of the decomposition of salicine, who ascertained its nature and composition.

5322. To obtain it, the oil of spirma is distilled with an aqueous solution of potash in excess as long as any oil distils. The residue, a solution of saliculite of potash, is supersaturated with dilute sulphuric acid, and again distilled, when saliculous acid comes over with the vapour of water. Or, according to Piria, a mixture of one part of salicine, one part bichromate of potash, two and a half of oil of vitriol, and twenty of water, is to be subjected to distillation. The salicine being dissolved in part of the water, and the acid diluted with the rest, the whole materials are mixed in a retort, when heat is excited with a gentle effervescence. When this ceases, the distillation should be commenced. Half a pound of salicine yields about two punces of saliculous acid. In both processes the distilled liquid contains saliculous acid, which separates from the water. It is purified by washing with water and rectification with chloride of calcium.

1.1731, which boils at 370°, or, according to Piria, at 380°, and congeals at — 4°. It has a burning taste, and a pleasant aromatic odour; is somewhat soluble in water, and mixes, in all proportions, with alcohol and ether. Its solution first reddens, then bleaches litmus. It is decomposed by concentrated sulphuric acid. When placed in contact with chlorine or bromine, one eq. hydrogen is removed, which, with those elements severally, forms chlorohydric or bromohydric acids; and is replaced by one eq. chlorine or bromine, producing chlorosaliculic, or bromosaliculic acid. Saliculous acid, treated with an excess of hydrate of potash, evolves hydrogen

gas, while saliculic acid is formed. Saliculous acid likewise decagagety drogen by reaction with potassium, forming saliculite of potast.

tallic oxides, its basic water being replaced by one eq. metallic oxides, its basic water being replaced by one eq. metallic oxides resulting fixed alkaline and ammoniacal saliculities are soluble, and of an alkaline reaction. The rest are insoluble. Most of them are low, and contain water of crystallization. A solution of the account the salts of peroxide of iron of an evanescent purple colour. In water copper it produces a green precipitate. Saliculous acid is separate to the salts of peroxide of iron of an evanescent purple colour.

the saliculites by the stronger acids.

mass, is prepared by adding concentrated liquid ammonia to salcolous. It is without taste, having a faint odour of roses; sparingly soluble out change of colour, in cold water and alcohol; more readily described hot alcohol. By the cooling of a saturated solution, transparent pakent needles are obtained. It is spontaneously decomposed if kept must; coming black, semi-fluid, and exhaling ammonia with a penetrating materials. Dry saliculous acid readily absorbs dry ammonia; and the pound, according to Ettling, contains three atoms of the acid and two of ammonia.

three of alcohol, and ammonia added by successive drops, the liquid cretes into a solid mass of fine yellow needles. With the aid of a mode heat these crystals dissolve; and the solution, by repose, deposits go yellow, brilliant, transparent prisms, which, when dry, are hard and verizable. Here three atoms of the acid (or one atom considered as basic acid) are acted on by two of ammonia, six atoms of water being nated. The alcoholic liquid in which these crystals have been formation to longer able, even at a boiling heat, to dissolve them. They now to a threefold quantity of alcohol. This would indicate, that at first solid of ammonia is formed, which, by a longer contact with ammonia, allow separation of the crystals, passes into saliculamide. This body it composed by a high temperature. Heated with acids and alkalies, it is solved into saliculous acid and ammonia. It is insoluble in water. It mula, C48 His O6 N2.

5327. Saliculite of Potash; Neutral.—This is best obtained by add saliculous acid to a warm solution of potash in alcohol, and allowing whole to cool, when the salt separates almost pure in the form of four-in pearly tables, nearly colourless, very soluble in water, spontaneously decaposed by exposure to the air in a moist state. It contains water of crystation, which is expelled by a heat of 212°. Formula, 2(C<sup>14</sup> H<sup>5</sup> O<sup>2</sup>) + KO

If the neutral salt be dissolved in hot alcohol, and saliculous acid added, acid salt is deposited, on cooling, in yellowish-white, long, fine, and brill needles. When dry it becomes yellow at a temperature of 230°. We decomposes it into a neutral salt, and saliculous acid, which separates.

5328. Saliculites of Soda, Lime, Baryta, and Magnesia, may formed directly. They have the properties of the potash salt. The sal soda contains two atoms of water of crystallization, removable by a han 230°. There is likewise an acid salt of soda in fine shining needles. I liculite of copper is anhydrous and green. The salts of zinc and mass are yellow and insoluble.

5329. Basic Saliculite of Lead.—Saliculous acid being dissolved dilute alcohol, and acetate of lead added to the boiling solution, on cooli

deposited, and may be separated by boiling alcohol from any adid. It is a lemon-yellow powder, which, when heated, froths up, water and acid; insoluble in water. Its formula is C<sup>14</sup> H<sup>5</sup> O<sup>3</sup> + If saliculous acid be added to diacetate of lead, a yellow powder of composition is precipitated.

Saliculite of Silver.—A solution of nitrate of silver causes, with iculite of potash, a greenish-yellow precipitate, which, when heated, d without disengagement of gas, the vessel being silvered by the

netal.

Melanic Acid, C<sup>10</sup> H<sup>4</sup> O<sup>5</sup>.—Discovered by Piria. When salicutash is exposed to the air, it acquires a green colour, which, after e, becomes black. When no further change is perceived, the sacid is completely converted into acetic acid and a black powder c acid thus formed is in the exact proportion to combine with all h contained in the original salt. The black powder possesses acid, and unites with bases; from which circumstance it has received

Three equivalents of oxygen and two of water unite with one atom lite of potash, and convert its acid into one equivalent of acetic,

squivalent of melanic acid.

Saliculic Acid, C<sup>14</sup> H<sup>5</sup> O<sup>5</sup> + HO.—Discovered by Piria. This merated by heating saliculous acid with caustic potash. The mix-st assumes a brown colour, but the heat must be continued until it white. At this time hydrogen is disengaged. The residue is to ed in water and treated with a mineral acid, which separates the icid. In order to obtain it pure it must be repeatedly crystallized. acid is likewise formed, when coumarin (stearopten of the Tonka reated with potash in a similar manner.

Saliculic acid sublimes without decomposition, and may be thus in the form of long crystalline needles, very similar in their apto benzoic acid. It crystallizes from water in fine tufts. This places with difficulty in cold water, but very easily both in hot I in alcohol. Sulphuric acid decomposes it when they are heated

The salts of this acid have been little examined. The saliculate s insoluble in water.

Chlorosaliculic Acid, also called chloride of salicyl, chloride of Dry chlorine gas is passed through anhydrous saliculous acid as hlorohydric acid is disengaged. On cooling, the compound belid and crystalline. It may be purified by crystallization from a plic solution, which deposits it, on cooling, in the form of pale yelue, rhombic tables, of a pearly lustre, having a peculiar aromatic d the capability of being sublimed without alteration. It is inflaming with a green flame, is insoluble in water, but soluble in alcother. It combines with alkalies, and may be separated from them d by acids. An alcoholic solution gives with acetate of copper a yellow, and with acetate of lead a yellow precipitate. Persalts of tinged by it of a dark blue. When heated with potassium, it is ed with heat and light. Ammoniacal gas converts it into chlorode.

This chloracid is distinguished from all analogous compounds of with compound radicals, by its power of combining with bases, and ag the action of those bodies. It forms with metallic oxides pecu-

liar salts, in which one atom of chlorosaliculic acid is c thined with atom of metallic oxide, supposing the oxygen and chlorine to change place that they may be considered as compounds of saliculic acid with achieves, C<sup>14</sup> H<sup>5</sup> O<sup>5</sup> + MCl, like the compounds of the chlorosaliculic acid with alkalies or metallic chlorides. Formula of chlorosaliculic acid H<sup>5</sup> O<sup>5</sup> Cl.

5339. It hence appears to be saliculimide, in which three atoms of

drogen are replaced by three atoms of chlorine.

5340. Bromosaliculic Acid.—This compound, in its preparation, perties, and action with ammonia, is completely analogous to the

ceding.

5341. Iodosaliculic Acid.—Saliculous acid dissolves iodine in quantity, without apparent decomposition. But iodosaliculic acid mobilized by distilling iodide of potassium with chlorosaliculic acid. I blimes in the form of a dark-brown fusible mass, analogous in its religious.

to the two preceding compounds.

with moderately strong nitric acid, is converted, with disengagement of a trous acid, into a crystalline mass of nitrosaliculic acid, which is public by washing with water, solution in alcohol, and crystallization. By state taneous evaporation, the alcoholic solution yields small transparent public the skin and nails permanently yellow, precipitates the salts of lead yellow, and those of copper green. It is inodorous, but has an acrid taste, excise cough. Heated with potassium, it is decomposed with explosive ignition. It combines with alkalies to form crystallizable compounds, which detonate when dried and heated. Ammonia colours the acid a deep blood-relation of iron is coloured cherry-red by it. These compounds demand a more accurate study.

5343. Furning nitric acid acts violently on saliculous acid, producing a volatile yellow matter, and a fixed residue containing a crystallizable stance not yet examined.—Gregory's Translation from Liebig.

# Of the Acids from the Oil of Gaultheria.\*

5344. From the observations and experiments of Mr. Procter, Jr., published in the American Journal of Phar-

<sup>&</sup>quot;Oil of Gaultheria Procumbens.—This volatile oil is extensively used by the pharmaceutists of this country to flavour syrups, etc. Most of the oil used in this city is obtained from distillers residing in New Jersey, in which State the plast yielding it grows in great abundance. As usually found in the shops, it has a most or less intense red colour; but when recently distilled it is colourless, or nearly to density, as the result of several careful observations, is 1.173, and its boiling point 412°, Fahr.; the mercury remaining stationary at that point. Its taste is burning and aromatic; it is slightly soluble in water, to which it communicates its odograms taste; and it mixes with alcohol and other in all proportions."—Procter, page 212.

reat resemblance between the habitudes of oil of gaulneria, and that from spiræa ulmaria, described above as aliculous acid. It appears either that salycyl exists in ne oil of gaultheria, and is productive of compounds reembling those produced by like reactions with the oil of pirea ulmaria; or that another compound radical exists the oil of gaultheria, which has a close analogy in proerties to salycyl.

5345. I will here quote the account given by Mr. Procer, Jr., in which the habitudes of the two oils in question

re contrasted.

5346. Mr. Procter premises in the following words:—"For several ears past it has been supposed\* that the volatile oil of the gaultheria proumbens, either from the analogy of their odour or specific gravity, possed similar properties with the oil of spiræa ulmaria, without any steps awing been taken to ascertain the correctness of the supposition. The observations which follow are intended to throw light on this subject. The bernical characteristics of oil of gaultheria have been found, in many insures, to accord with those described as peculiar to saliculous acid, yet everal instances occur to the contrary.

5347. "They have the same density, and the aqueous solution of each olours the persalts of iron purple. The compounds which potassa, soda, and oxide of copper form with oil of gaultheria, are very like the salts of

aliculous acid with those bases.

5348. "The action of an excess of caustic potassa with heat produces a rystalline body, identical in all its reactions with saliculic acid, as described y Piria.

5349. "The compound of oil of gaultheria and potassa, when exposed to be combined influence of moisture and the atmosphere, undergoes a decom-

osition similar to that of saliculite of potassa.

5350. "The reactions of chlorine and bromine with oil of gaultheria ield compounds similar to those with saliculous acid; and nitric acid also

roduces results of an analogous character.

5351. "On the contrary, the boiling point of oil of gaultheria is many egrees higher than that of saliculous acid. Ammonia forms a compound rith it which differs from saliculite of ammonia in not being decomposed by cids with the separation of the oil, nor by potassa with the separation of momonia. All endeavours to form the body called saliculimide by Liebig, with the process he gives, were ineffectual. The compounds of baryta and ead with oil of gaultheria are white, while the saliculites of those bases are rellow. But the most striking difference between these substances is, that when oil of gaultheria is boiled with solution of potassa, it is not recoverate by means of an acid, as saliculous acid is. Under these circumstances a crystalline substance is precipitated, which is the same acid that results from heating the oil with an excess of potassa."

5853. "Dropped into a concentrated soluti of or sode, their is instantly solidified, becomes white, and separates from the alkalist shiften while heat is disengaged.

5353. "Oil of gaultheria decomposes the carbonaiss of potassa and misgradually, without heat; but if gently warmed, the evolution of entires

acid is syident.

5354. "Chlorine and bromine, when brought into contact with old gaultheria, combine with it; the mixture becomes very hot, and hydrobine ric and hydrobromic acids are evolved. Indine is dissolved by the old forming a deep red solution without combining with it, as heat display the indine without the production of any hydriodic acid.

5355. "Nitric acid of density 1.40, assisted by heat, converts oil of pattheria into a crystalline substance having acid properties, whilst nitrous of fumes are evolved. If fuming nitric acid be employed, the reaction is the lent, without the assistance of heat, and a different product is obtained.

5856. "When oil of gaultheria is added to concentrated sulphuric acid, the latter becomes elightly coloured, and, if heated, the odour of the side

destroyed.

5367. "When oil of gaultheria is distilled with solution of putasse in the case, the distilled liquid has neither the odour nor tasts of the oil, and as sequently its constitution differs from that of the oil of spirms ulmaria, which under the same circumstances, yields a volatile oil distinct from salicular soid, that acid remaining combined with the potassa."

5358. For further particulars respecting the results of Mr. Procter's meritorious investigation, I refer to his memoir. I hope that the brilliant success which has attended his efforts, may cause them to be emulated by many of his countrymen.

#### OF URIL.

#### Or Cyanoxalic Acid.

bination of carbon, nitrogen, and oxygen, of which the formula is C<sup>n</sup> N<sup>n</sup>O<sup>n</sup>, and which may be considered either as a compound of carbonic acid acid cyanogen, or as resulting from the substitution of cyanogen for one of the three atoms of oxygen in oxalic acid. Thus the formula of the latter beat C<sup>n</sup> O<sup>n</sup>, that of uril will be C<sup>n</sup> O<sup>n</sup>+Cy. It may be remembered that ures, a crystallizable matter of the urine, has been mentioned as being equivalent eyanate of ammonia, C<sup>n</sup> NO + NH<sup>n</sup>O, or more properly a cyanate of a oxide of ammonium, this oxide comprising the elements of one atom of monia, NH<sup>n</sup>, and one of water, HO (1307).

5860. This being premised, the following formulæ of the combinations

uril will, it is presumed, be understood.

2 atoms uril +  $\begin{cases} 1 \text{ urea} = 1 \text{ uric acid, } C^{10} \text{ N}^4 \text{ H}^4 \text{ O}^4 \\ O^4 + 4 \text{HO} = \text{alloxan, } C^6 \text{ N}^6 \text{ H}^4 \text{ O}^{10} \\ O + 5 \text{HO} = \text{alloxantine, } C^6 \text{ N}^6 \text{ H}^4 \text{ O}^{10} \\ 1 \text{ ammonia} + 2 \text{HO} = \text{uramile } C^6 \text{ N}^6 \text{ H}^5 \text{ O}^6 \end{cases}$ 

# If Uric Acid, and various Substances to which it gives rise.

#### Of Uric Acid, C10 H4 O6 N4.

5361. As the most frequent and abundant material in rinary calculi, uric acid, and the substances which conribute to its formation, or which may be derived from it, perit the most sedulous attention.

5362. This acid is an ingredient in the urine of men, and generally in that of carnivorous animals, forming, as lready mentioned, calculi, depositions from urine, and youty or arthritic concretions. In the state of urate of immonia, it constitutes the greater proportion of the extrement of the boa constrictor and other serpents; also of pirds, more especially those of the carnivorous species. I bound it to abound in that of a young eagle. An accumutation of the excrement of certain aquatic birds, containing large amount of this acid, on some islands near the coast of Peru and Chili, under the name of guano, is much used as manure.

5363. I infer that the best process for obtaining uric acid is as follows:—Boil the substance from which it is to be extracted in a dilute solution of caustic potash. On allowing the decoction to cool, urate of potash, which is almost insoluble in cold water, precipitates, leaving some impurities in solution. After being well washed with water, the urate thus separated is redissolved in a boiling solution of potash, next filtered, while hot, and afterwards added to an excess of chlorohydric acid maintained in a state of ebullition. The uric acid which precipitates, is to be rendered pure and white by repeated aqueous ablution.

5364. Liebig recommends, that in extracting uric acid from excrement, a solution of borax be employed, as it does not take up so large a portion of the impurities as caustic potash.

5365. Uric acid crystallizes in thin spangles, with a dazzling white satin lustre. It is insipid and inodorous. At a boiling heat the crystals sustain no loss of water. It is heavier than water, almost insoluble in that liquid when cold, and but little soluble in it when hot. The solution feebly reddens litmus.

5366. When to a well refrigerated aqueous solution of

borate of soda, holding uric acid dissolved, chlorohydiacid is added, the uric acid precipitates in a hydrau state, forming a transparent jelly, which, by a feeble bet, is converted into a crystalline powder, consisting of anydrous uric acid.

5367. This acid is soluble in concentrated sulphuric acid, but separates on dilution with water. It is not soluble in concentrated chlorohydric acid than is pure water.

5368. Subjected to dry distillation, it yields the same products as urea, that is to say, cyanic acid, cyamelit, cyanhydric acid, a little carbonate of ammonia, and a brown and carbonaceous residue very rich in nitroges. During this decomposition, the hydrated cyanic acid and ammonia unite in the neck of the retort, forming urea.

5369. In dilute nitric acid, uric acid dissolves, with lively effervescence, from the escape of equal volumes of carbonic acid and nitrogen. The resulting solution contains alloxan, alloxatin, urea, parabanic acid, and analogie.

5370. By the addition of an excess of ammonia, the concentrated liquid becomes purple red, from the generation of murexide. This effect is one of the means of recognising the acid.

5371. Fused with hydrate of potash, uric acid produces, with the alkali or its metal, a carbonate, a cyanate, and a cyanide.

5372. Subjected to boiling water with the bioxide of lead, it is resolved into allantoin, oxalic acid, and ures. Heated to 320°, with a little water in a tube hermetically sealed, this acid is dissolved, without the evolution of gas forming a yellow transparent liquid, which, on lowering the

temperature, assumes a gelatinous appearance.

5373. Uric acid is peculiar in combining with metalic oxides, without abandoning water. The urates of the alkalies and alkaline earths are little soluble in cold water, but very soluble in this liquid when boiling, the solubility being augmented by an excess of alkali. The urates formed with other metallic oxides and with ammonia, are white and insoluble. All the urates are easily decomposable by acids, even by the acetic acid. When first liberated, uric acid assumes the form of a jelly, which is soon changed into fine brilliant spangles.

5374. Allantoin is a substance arising from the urine of a fetus in the exterus of a cow, and may be obtained from the waters of the allantois of this animal, by evaporation and crystallization. It may be more easily procured by the following means:—To uric acid, diffused through twenty parts of boiling water, freshly prepared, bioxide of lead is to be added as long as the colour is affected. The boiling liquor is to be filtered, and evaporated until crystallization commences. It is then allowed to cool in a quiescent state. By this procedure the allantoin separates in crystals, an covalate of the protoxide of lead being simultaneously produced.

5375. In order to understand this process, the composition of the materials and products should be remembered. They are as follows:—Uric acid, C10 H4 N4 O6; bioxide of lead, one atom of protoxide, one of oxygen;

area, C<sup>3</sup> N<sup>3</sup> H<sup>4</sup> O<sup>3</sup>; allantoin, C<sup>4</sup> N<sup>4</sup> H<sup>5</sup> O<sup>5</sup>; oxalic acid, C<sup>3</sup> O<sup>3</sup>.

5376. Hence, assuming that five atoms of water are taken up, and one atom of oxygen for each of the four atoms of bioxide of lead, the materials are:-

| Two atoms of uric acid, - Five water, Four oxygen,                       | •             | • | Cao | H *<br>H * | N° | O18<br>O 5<br>O 4 |
|--------------------------------------------------------------------------|---------------|---|-----|------------|----|-------------------|
|                                                                          |               |   | C20 | H18        | N° | O <sup>91</sup>   |
| The products are two urea, Four oxalic acid in the oxala: One allantoin, | -<br>te,<br>- | • | C · | H *        |    | 018               |
|                                                                          |               |   | C20 | H18        | N° | Ozi               |

The protoxide of lead being in both of the aggregates, does not affect the

result by being omitted from both.

5377. It is from the results of this reaction between uric acid and bioxide of lead, that Liebig has inferred the existence of uril as above mentioned (5319).

5378. Alloxan, or erythric acid, is one of the products which have resulted from the decomposition of uric acid. To prepare alloxan, one part of uric acid is to be added to four of nitric acid, of a density between 1.41 and 1.5. As the reaction causes much heat and effervescence, the uric acid should be added in successive small portions. Little white granular brilliant crystals are gradually formed, until the whole becomes one aggregate of them, which, after being allowed to drain in a glass funnel, must be dried on porous brick, of porcelain earth. By re-solution and re-crystallization,

the crystals of alloxan, thus formed, will be rendered quite pure.

5379. Alloxan crystallizes in octahedra, with rhomboidal bases, colourless, transparent, very brilliant, and often of an inch in diameter. They are efflorescent, losing 25 per cent. of water. By a gentle heat, alloxan, thus crystallized, is rendered anhydrous. It may be obtained in anhydrous crystals, in the form of oblique rhomboidal prisms, which resemble rhomboidal octahedra with truncated summits, from an aqueous solution of allozane saturated while hot. It is very soluble in water, has a nauseous smell, and a salt and feebly astringent taste. It reddens vegetable colours, and tinges the skin purple. By reaction with alkalies, it is decomposed into alloxanic acid. Boiled with an alkali, it is transformed into urea and meschloride of tin, or metallic zinc, and chloronyoric acid. An excess of meanic acid; sulphuric or chlorohydric acid into allocantin; sulphurous and ammonia into thionurate of ammonia; allocantin and ammonia of iron, it produces an indigo-blue liquid. With metallic cardes a cambine without decomposition.

by the metamorphosis of alloxan by caustic alkalies. The anadromical contains the elements of half an atom of alloxan, minus one atom of water.

2HO, is one of the products of a solution of alloxanate of baryla of states, saturated at a boiling heat. Also, when a solution of alloxanate parties, drop by drop, into a boiling solution of a setate of lead, a very heavy frame har mesoxulate of lead precipitates, while nothing remains in the seed lead besides the excess of acetate of lead and pure urea. Both this and the proceeding acid may be separated and crystallized. They are powerful solutions.

5862. Mycomelinic acid, C<sup>16</sup> N<sup>o</sup> H<sup>10</sup> O<sup>10</sup>, is formed on adding an cool ammonia to a solution of alloxan, and raising the mixture to the big point. It is almost insoluble in cold water, and is thrown down as a low gelatinous precipitate, which becomes a yellow porous powder on the

5888. Parabasia said, C\* N\* O\* + 2HO, is one of the products of decomposition of uric acid or alloxan by nitric acid, discovered by Line and Woshler. It is prepared by dissolving one part either of uric acid alloxan in eight parts of nitric acid of ordinary strength, evaporating in liquor to a syrup, and allowing it to crystallize.

5384. It has a very sour taste, resembling that of oxalic acid, and for thin transparent six-sided prismatic crystals. It is very soluble in the

and does not effloresce; it is in some degree volatile.

5365. Oxuluric acid, C<sup>o</sup> N<sup>o</sup> H<sup>o</sup> O<sup>o</sup> + HO, is formed on adding exacts to a boiling solution of parabanic acid, or on supersaturating with sames a solution, recently prepared, of uric acid in nitric acid, which yields, be evaporation, crystals of oxalurate of ammonia. The acid, when separate is a light brilliant white crystalline powder; its taste is very sour, and reddens litmus. Its aqueous solution is decomposed completely by stall tion, and resolved into oxalic acid and oxalate of urea. It is formed by the combination of the elements of parabanic acid with two atoms of wall The crystallized acid contains the elements of two atoms of oxalic acid so of one atom of urea, and may be considered as uric acid in which the sill is replaced by oxalic acid.

5386. Thiomuric acid, C\* N\* H\* O\* (S\* O\*) + HO, is a bibasic so produced by the simultaneous action of sulphurous acid and ammonia ax alloxan. Liberated from thionurate of lead by sulphuretted bydrogm, crystallizes in very thin needles, is persistent in air, very soluble in wate and has an acid taste. It contains the elements of one atom of alloxan, at

Finding Graham's Elements to contain an abridgment of the account given! Liebig, of the compounds, or products, of uric acid, I have made a free use of it, we such changes in the language as to make it my own, where it was not such a should have used. In some cases I have made a similar use of Kane's Element and of Gregory's Translations from Liebig, especially in the account of saliculated and its compounds.

nonia, and two atoms of sulphurous acid. On heating thionuric ms of oxygen of the alloxan reunite with two atoms of sulto form sulphuric acid, while the elements of urile, ammonia, ombine and give rise to uramile.

amile, C<sup>8</sup> N<sup>3</sup> H<sup>5</sup> O<sup>6</sup>, is prepared by adding hydrochloric acid d and boiling solution of thionurate of ammonia, till it is strongly heat is continued till the liquid begins to become turbid; it is to cool for crystallization. Uramile crystallizes in thin and or presents itself in the form of a brilliant white powder comy thin silky needles. It is sparingly soluble in hot water, able in cold water, dissolves in ammonia and caustic alkalies, precipitated, without alteration, by acids. Either diluted acids of potash, boiled upon uramile, convert it into uramilic acid, ammonia. The ammoniacal solution of uramile becomes purand deposits crystalline needles of a green colour and metaln contact with oxide of mercury or oxide of silver, it is decomullition, into murexide, and at the same time reduces the oxides ic state.

amilic acid, C<sup>16</sup> N<sup>5</sup> H<sup>10</sup> O<sup>15</sup>, is prepared by dissolving thionuonia in cold water, adding to the saturated solution a small
ulphuric acid, and evaporating by a water-bath. After a time,
l is deposited in transparent four-sided prisms of a vitreous lusty needles. It is soluble in six or eight parts of cold water,
parts of boiling water, forming a feebly acid solution. For the
tramilic acid, two atoms of uramile unite with the elements of
of water, yielding up, at the same time, the elements of one
nonia.

loxantin.—Formula, C<sup>8</sup> N<sup>9</sup> H<sup>5</sup> O<sup>10</sup>. Alloxantin was first obr. Prout among the products of the decomposition of uric acid i, and more lately produced and studied by MM. Liebig and whom several processes are given for its preparation. 1. From One part of uric acid is boiled with thirty-two parts of water, tric acid added, by small portions at a time, till the uric acid is issolved, and the liquor evaporated to two-thirds. In the course s, or sometimes a few hours, the alloxantin is deposited in crysre purified by new crystallizations. 2. From alloxan.—It is large quantity by conveying a stream of sulphuretted hydrogen Sulphur is first deposited, and then the whole nick mass of crystals of alloxantin, which are separated from olution in boiling water. The alloxantin crystallizes by evapotate of purity. 3. On exposing a solution of alloxan to the acoltaic battery, oxygen is evolved at the anode, and alloxantin is the cathode in crystalline crusts.

loxantin crystallizes in oblique prisms of four sides, which are slightly yellow, hard, and easily reduced to powder; they beair impregnated with ammonia, and acquire a green metallic sy are not altered at 212°, but at 302° (150° centig.) lose three ter; are sparingly soluble in cold water, more soluble in boiling solution reddens litmus. Alloxantin heated in chlorine-water, nitric acid, is changed into alloxan. With salts of silver it prok precipitate of metallic silver. It is decomposed by alkalies; r produces, in its solution, a violet precipitate, which is made y heat, and finally disappears. By the action of boiling sul-

pheric acid, two atoms of alloxan are converted, t community two atoms of water, into one atom of alloxantin, three atoms of ammonia, and two atoms of carbonic acid.

plained by M. Liebig. By the action of nitric acid, the uril of the manifestation of mitric acid, the uril of the manifestation of mitric acid, the uril of the manifestation of water, giving rise to one atom of alloxantin, and to quadroxide of mitrigen, NO\*, which, in contact with water, is converted into nitrous and now acids; the nitrous acid is decomposed, and half of the urea set at heavy while the other half of the urea forms, with nitric acid, nitrate of urea to the process with sulphuretted hydrogen, one atom of oxygen of the allows combines with hydrogen from the sulphuretted hydrogen to form war, which remains in the constitution of the alloxantin; the sulphur set from

deposited.

5892. Products of the decomposition of Allexantin. - When a stru of sulphuretted hydrogen is carried into a boiling solution of allowants more sulphur is deposited, and on saturating the solution with anymons, salt crystallizes in thin colourless needles, of which the formula is C N H<sup>7</sup> O<sup>a</sup>, which is considered a compound of a new acid, disturic acid, with ammonia. This acid is resolved into new products when liberated by each ther acid; one of these produced by exposure to air, and evaporation of the solution of the ammoniacal salt in dilute sulphuric or hydrochionic acid. dimorphous allocation, a body having the same composition as allocation with a different form. On mingling boiling solutions of sal agranguise and alloxantin, the mixture becomes suddenly of a purple-red colour, then go dually loses its colour, becoming turbid, and deposits colouries imitself plates of uramile, which become rose-red on drying. The liquid costs after its decomposition, alloxan and free hydrochloric acid. tion of alloxantin is heated with caustic ammonia, uramile and mycomesnate of ammonia are first formed, but are decomposed into other pushed by the prolonged action of ammonia and air. A recent solution of allessetin in ammonia gradually absorbs oxygen from the air, and deposits caytals of exalurate of ammonia.

#### Murexide.

5393. Formula, C<sup>13</sup> N<sup>5</sup> H<sup>6</sup> O<sup>3</sup> (Liebig and Weehler). This beautiful product of the decomposition of uric acid was first described by Dr. Prost under the name of purpurate of ammonia. Murexide may be formed by evaporating a solution of uric acid in dilute nitric acid, until it acquares flesh-red colour, and treating it, when cooled to 160°, with a dilute solution of ammonia, till the presence of free ammonia is perceptible; the liquid in the diluted with half its volume of water, and allowed to cool. It may also be formed by the contact of ammonia with various other products of the reaction of nitric acid with uric acid, with ammonia, with or without the presence of atmospheric air.

5394. The following method, proposed by Liebig and slightly modified by Gregory, appears to be the easiest and most certain, and also most productive:—Seven grains of hydrated alloxan, and four grains of alloxasts, are dissolved by boiling in 240 grains of water, and the boiling solution added to 80 grains, by measure, of a cold and strong solution of carbonal of ammonia. This mixture has precisely the proper temperature, and posits very fine crystals of murexide. The experiment is not so successful.

a large scale; probably because the liquid, by remaining longer warm, dergoes a partial change. It is best to try first a saturated solution of rbonate of ammonia in cold water. If it do not yield good crystals, add little water, and repeat the experiment till a solution of the carbonate is tained, which gives a good result. The difficulty is owing to the spontature of the shops; but when a proper solution is obtained, the experiment wer fails.

5395. Murexide crystallizes in short four-sided prisms, of which two zes, like the upper wings of cantharides, reflect a green metallic lustre. crystals are garnet-red by transmitted light. Their powder is reddishown, and acquires a green lustre under the burnisher. Murexide is but ghtly soluble in cold water, but colours it of a magnificent purple; it dis-Ives, however, readily in water at 158°, and crystallizes again as the sotion cools. It is insoluble in alcohol, ether, or in water saturated with rbonate of ammonia. But this substance cannot be purified or obtained crystals of large size by crystallizing it from boiling water; for on boilg murexide in a small quantity of water for the time necessary to dissolve s whole, the crystals become colourless, and, upon cooling, a yellow gelanous matter precipitates. Hence, probably, the slight uncertainty which tends even the best process for the preparation of this substance. e dissolves in a solution of potash, producing a superb indigo-blue colour, hich disappears with the application of heat, ammonia being disengaged. If the inorganic acids decompose murexide, precipitating from its solution urexan in small brilliant plates. Sulphuretted hydrogen decomposes it amediately into alloxantin, dialuric acid and murexan, while sulphur is set

5396. Murexan, C<sup>6</sup> N<sup>9</sup> H<sup>4</sup> O<sup>5</sup>, was named purpuric acid by Prout. It formed on dissolving murexide with heat in caustic potash, heating till be blue colour disappears, and then adding an excess of dilute sulphuric id. It crystallizes in colourless plates which have a silky lustre, and are ray brilliant; is insoluble in water and dilute acids; it dissolves in ammoia and other alkalies, in the cold, without neutralizing them. The properess of murexan closely resemble those of uramile. Like uramile, murexan oiled with water, red oxide of mercury, and a little ammonia, yields murkide. The composition of murexan and uramile, also, not differing much 100 parts, Dr. Gregory admits it to be possible that these two substances may be essentially the same.

5397. As the habitudes of uric acid, and of the substances from which it my be generated, or to which it may give rise, must be an object of intest to the surgeon and physician, I have deemed it proper to make a colous abstract respecting it from Graham. I do not, however, as respects ther bodies, deem it expedient to go farther into these boundless regions of hemistry. The multiplication of compounds, rendered distinguishable in seir properties by shifting the associations of ponderable, with imponderable satter, seems to be as unlimited as the images which may be produced in se kaleidoscope, by varying the relative positions of the coloured beads: nd as, in a majority of instances, the compounds created by the changes lluded to, have either the electro-positive, or electro-negative character, thich distinguishes acids and bases from other bodies; so it must happen sat there will be a prodigious and increasing number of substances stumped ith the attributes of acidity or basidity. Even adepts in the science will

find it impossible to retain any available knowledge of the details response
such compounds, and of course, however important it may be to region that is known of them in systematic works, in a text book it can answer good purpose to dwell on that which could not be remembered even it were once well learned.

5398. I propose, however, in an appendix, to give some alphabeted to bles, in which the information, with which it were inexpedient to dog to

body of this work, may be found-

# On the Influence of Benzoic Acid in lessening the Generalist of Uric Acid in Human Urine.

5399. Allusion has been made to the discovery, by Alexander Ure, that benzoic acid, taken into the human stomach, is converted into hippuric acid, causing a dimme tion of the uric acid generated in the urine. The observe tions and inferences of Ure have been confirmed by the of Bouchardat, who alleges, that in the case of a patient the hospital of Hotel Dieu, at Paris, labouring under acti rheumatism, and whose urine was depositing an abundant of uric acid, the spontaneous deposition of the acid ceast after the due administration of benzoic acid: also, it is a leged by Mr. Garrod, that having repeatedly performed Ure's experiment, by swallowing from a scruple to half drachm of benzoic acid at a time, he had always been cutbled to obtain from his urine, passed three or four hours subsequently, on the addition of hydrochloric acid, from fifteen to twenty-nine grains of hippuric acid.

5400. There is, however, the opposite testimony of a commission of the French Academy of Sciences, drawn up by Gay Lussac and Pelouze, that they could not find any verification of the results of Mr. Ure. Agreeably to the knowledge which I have obtained respecting the manual in which such commissions are managed by some of the most distinguished of the academicians, I attach very little importance to their negative testimony. With excellent intentions, they are too much occupied, too much distinguished of the academic and the intentions.

tracted, to do their duty well in such cases.

5401. I have not met with any statement tending to explain in what manner the elements of benzoic and uric acid can give rise to hippuric acid.

Bell's Pharmacoutical Journal, London, page 50. No. 12. June, 1862.

#### OF ORGANIC ALKALIES OR BASES,

alled Vegetable Alkalies, Vegeto-Alkalies, or Alkaloids.

2. The discovery of the substances which bear the mentioned names, is of the highest importance to nd. It has enabled the physician to avail himself of tive principles of some of the most powerful remerith a certainty which was before unattainable. The t, in lieu of being nauseated and even injured by of which the greater part, perhaps the whole, may rt, if not injurious, has to swallow nothing which

inefficacious, when judiciously prescribed.

- 3. The organic alkalies are entitled to rank as bases, the definition of acidity, deduced from the practice nists, and given in this work (note 631), that whatiturates a well defined acid must be deemed a base. 1. The compounds formed with acids by the alkaline under consideration, resemble those formed with c oxybases; their acids and ingredients being no less tible of precipitation by the appropriate tests. Thus ulphates are liable to be deprived of their acids by a n of baryta, their chlorides by solutions of silver or There is in this respect a striking difference between bitudes of these organic alkaline bases and those are formed of oxidized compound radicals, like the of ethyl, formyl, and methyl, which cannot be transfrom one acid to another, unless in a nascent state, er peculiar circumstances. Even when isolated, the last mentioned refuse to unite with hydrated acids, is far from being the case with the alkaline bases in Generally, the latter differ very much from the s proper, in being much more soluble in alcohol than er. In consequence of this last mentioned trait their e reaction with vegetable colours is very feeble, beplayed more in their power of restoring such colours, directly producing the changes which result from ns of the inorganic alkaline bases.
- i. The following table of the organic alkalies indiheir sources and composition:—

| NAMES.                           |            |                                   |                | 080    | BOURCES.      |          |   |     | COMPOSITION.*                                  |
|----------------------------------|------------|-----------------------------------|----------------|--------|---------------|----------|---|-----|------------------------------------------------|
| Morphiat or morphine,            |            | Opium,                            |                |        |               |          |   | ٠   | CarH18OoN                                      |
| Paramorphia or paramorphine.     | _          | Do.                               |                | ٠      | •             | ٠        | • | •   | CALLACA                                        |
| Pseudomorphia or pseudomorphine, | _          | Do.                               |                |        | •             | •        |   | •   | Callinown                                      |
| Codeia or codeine,               | _          | Do.                               |                |        | •             | ٠        |   | ٠   | C31H20ON                                       |
| Narcotina or narcotine,          | _          | Do.                               |                |        |               |          | ٠ | •   | Cao HanOus N                                   |
| Narceia or narceine,             | _          | Do.                               |                | •      |               | •        | ٠ | •   | Cas Hay OreN                                   |
| Cinchonia or cinchonine,         | 14         | Peruvian bark,                    | ark,           |        | •             |          | • | •   | CoH <sub>18</sub> ON                           |
| Quinia or quinine,               |            | Do                                |                |        |               |          | ٠ | ٠   | Co.HaOaN                                       |
| Aricina or aricine.              |            | Bark of an unknown tree,          | unkno          | wn tre | , g           | ٠        | * | ٠   | NOn Had                                        |
| Strychnia or strychnine,         | <i>G</i> 2 | St. Ignatius's bean, nux vomica,  | 3's bea        | n, nux | vomica,       | dec.     |   | ٠   | CaH <sub>1</sub> O <sub>5</sub> N              |
| Brucia or brucine,               |            | False Angu                        | ngustura bark, | bark,  |               |          |   | ٠   | C#H#ON                                         |
| Delphia or delphine,             | G/2        | Stavesacre,                       |                |        |               | •        |   | ٠   | Ca.HisOsN                                      |
| Verntria or veratrine,           |            | White hellebore                   | spore 8        | and me | and meadow as | saffron, | ٠ | *   | N <sub>2</sub> O <sub>2</sub> H <sub>2</sub> O |
| Sabadillia or sabadilline,       | -          | Cevadilla,                        |                |        |               |          |   | •   | Collia05N                                      |
| Emetia or emetine,               | _          | Ipecacuanha,                      | eg,            | •      | •             |          | • | ٠   | N.O.H.O                                        |
| Solania or solanine,             |            | Bittersweet and black nightshade, | and b          | lack n | ightshad      | 9        |   | •   | C*H*O*N                                        |
| Atropia or atropine,             |            | Belladonna,                       |                |        |               |          |   | •   | C*II*OHN                                       |
| Menispermia or menispermine,     |            | Cocculus Indicu                   | dicus,         |        | •             | •        | ٠ | •   | CASTANTA                                       |
| Melamia or melamine,             | -          | Artificial                        | ŀ              |        | . ,           |          |   | + 8 | CHICANO                                        |

| serving or jerving,             | ' A cramman man man man A l'                        | }          |
|---------------------------------|-----------------------------------------------------|------------|
| Conina or conine, or conicine,  | . Hemlock or conien maculatum,                      | CaHi       |
| Caffeina, caffeine, theine, or  | Curana, a paste made of the grain of paullinia      | CellsNans  |
| guaranine,                      | sorbilis, or from coffee, or tea,‡                  | ONUO       |
| Chelerythrina or chelerythrine, |                                                     |            |
| Colchicina or colchicine,       | . Colchicum atumnale,                               |            |
| Coradalina or coradaline,       | . Root of corydalis bulbosa and fabacca, .          |            |
| Curarina or curarine, .         | Curara poison,                                      | Formulæ    |
| Daturina or daturine,           | Datura stramonium,                                  | not given. |
| Cissampolina or cissampoline,   | Root of cissampelos pariera,                        | )          |
| Solanina or solanine,           | Several species of solanum; first shoots of potato, |            |

the quantity of each alkali in the anhydrous state, adequate to saturate one equivalent of an acid The several formulæ indicate

to young any saustactory course respecting the terminating monosyllable in the names of the organic alkalies. By it has been considered important to have the alkaline character signalized by the termination in a, while the first indebted for the discovery of the alkalies in question, and from whom, commercially, we have been in the Moreover it has not been pursued in some of the recent British compilations of the tained in neutral or indifferent principles. But this plan has not been adopted by the French and other continciples, have been variously estimated at different times; and it constantly happens that a principle previously he United States Dispensatory, and Pharmacoposia, the termination in a has been adopted, quinine and narcoting be treated as a base, agreeably to Bert by which the principle in question is In fact, the claim of this last mentioned principle to be considered as an alkali There is some difficulty in adopting any satisfactory course respecting the terminating monosyllable in the names of the organic alkalies. e circumstances, I have concluded to mention both names, but employ that by taken into the class of alkalies. Thus, for instance, caffein, or theine, is to rlance, than quinia or narcotina. for the purpose of the arts. highest rank. Hence, although in t termination in in or ine should be re some British and American chemist habit of receiving supplies of them are used much more, in common pe as well as those of several other pri considered as neutral, is afterwards Under the nental chemists, to whom we were # Berzelius' Report, 1841, 143. zelius' Report for 1841. usually known.

5406. The salts, formed by the organic alkalies with oxacids, always contain the elements of an atom of water essential to their existence. this respect they agree in their habitudes with the analogous ammonant compounds formed with the same acids. But in uniting with chlorohyan acid, or other halohydric acids," no water is requisite. In this respect this there is an agreement between their habitudes and those of a immonia. Here it might be reasonably inferred that in the one case the halogen body mass with a hydroret of the organic alkali, while in the other, the oxacid mass with an oxide of such a hydruret. This theory has made no change a the names of ammoniacal oxysalts; but as respects haloid compounds a ter changed muriate of ammonia into chloride of ammonium, and induced it analogous result in the case of the ammonineal compounds of each halogous body. Consistency then would seem to require that a like change should be made in the nomenclature of the compounds of the halohydra ach with the organic alkalies; but we have had no proof that any of those kalies are metallized, and of course could not call muriate of morphia chisride of morphium. Under these circumstances, chlorohydruret in the same to which I would resort for any compound of chlorohydric acid with an exganic base. In practice, however, until the relation between ammona and these alkalies is better understood, it will be as well to employ the officered appellation (muriate) above mentioned; keeping the other in view in order to prevent a theoretic misconception, that any combination can be formal with an organic base which merits to be designated as a muriate.

5407. The organic alkalies are, for the most part, products of vegetation; yet the following substances, not derived from vegetables, are alleged to be allied to the class of vegetable bases, ammeline, melamine, aniline, urea: also some substances obtained from the animal oil of Dippel, called severally odorine, ammoline, and animine.

### Organic Alkalies of doubtful Existence.

5408. "The following bases are still problematical: apirine, azaridine, blanchinine, buxine, carapine, castine, chioccine, crotonine, cynapine, daphnine, digitaline, esenbeckine, eupatorine, euphorbine, fumarine, glancine, glaucopicrine, jamaicine, menispermine, paramenispermine, pitayine, sanguinarine, staphisaine, surinamine, violine. Besides two bases in Carthagena quinquina bark and in chinova bark." Graham's Translation from Liebig, 983.

Of the State in which the Organic Alkalies exist in the Products of Vegetation, and the Means of extricating them, generally described.

5409. The organic alkalies appear in almost every instance to exist in the vegetables to which they belong, in

<sup>&</sup>quot;Halohydric is the generic name which I apply to acids formed of a halogen holy and hydrogen.

union with an acid. Thus, morphia is united with sulphuric and meconic acid, cinchonia and quinia with kinic acid, delphia with malic acid, and veratria with gallic acid. In some instances, the acids have not been specified; but the method requisite for the analysis, shows that they are present. The salt thus formed is entangled sometimes with resinous matter, sometimes with colouring matter, at others with fatty matter, and in a few instances with caoutchouc. In some cases several, in others all of these

impurities may be present.

5410. In the extrication of the organic alkalies characterized, and situated as has been stated, the first object of the chemist will be to employ some solvent which will take up the native salt in which it exists. This may in many cases be effected by water alone; but an aqueous solution of some powerful acid, usually sulphuric or chlorohydric acid, appears to have been found preserable. The next step is decomposition of the salt formed with the organic This may, of course, be effected by any stronger base, and accordingly, potash, soda, ammonia, lime, and magnesia, have all been more or less employed. The alkali when insoluble in water, as happens in a great majority of cases, precipitates with or without the precipitant, accordingly as the compound which this forms with the acid is or is not soluble. In either case, the next object to be attained is to extricate the organic alkali from the impurities which may have been precipitated with it. These may consist of resinous matter, fatty matter, colouring matter, caoutchouc, &c. To remove these, washing with weak alcohol, ether and water, has been employed, or re-solution in an acid, and subjection to the depurating and decolorizing efficacy of animal charcoal. Repeated solution and recrystallization by means of alcohol, or acids, are also used to effect a final depuration. When the alkali to be extricated is soluble in water, and volatile as in the instance of conicine, the leaves, flowers, roots or seed, are subjected, with a weak, aqueous, alkaline solution, to the distillatory process. The water which distils in consequence, contains more or less of the organic alkali, as well as some ammonia resulting from its decomposition. Being first neutralized by diluted sulphuric acid, then concentrated by evaporation, and afterwards digested in a close vessel with ether, this liquid dissolves the organic alkali,

which may of course be easily isolated by subsequent exposure to a water bath sufficiently heated to expel the ether and ammonia.

5411. In some instances the decomposition of the national salts in which the organic alkalies are constituents, may be effected by acetate of lead. As this metal generally forms insoluble compounds with vegetable acids, by complex affinity the acid goes to the oxide of that metal, while the alkali combines with acetic acid. From the solution of the acetate thus formed, the lead of any excess of the acetate of lead may be precipitated by sulphydric acid.

#### Of Morphia or Morphine.

5412. Morphia, the most important among the active principles of opium, was discovered by Serturner, of Embech, in Hanover, and recognised by him as an organical alkali. This formed the first step in a new career a chemical discovery, having induced those subsequent researches by other chemists, to which we are indebted for our knowledge of the series of analogous principles not tioned in the preceding table.

5413. Morphia exists in opium in chemical union with meconic acid only, but is mechanically associated with various substances, of which an account has been given (5172.)

5414. It is remarkable, that since we have learned the existence of morphia, it has become evident that the means of detecting its presence in laudanum, almost extemperaneously, had long been at hand in the shop of every druggist. Dr. Staples, a graduate of our school, demonstrated, about twenty years ago, that to cause a precipitation of crystals of morphia, it were only requisite to add to the tincture equal parts of liquid ammonia and alcohol. The crystals thus obtained, being redissolved by acetic acid.

\* The following process for elaborating the organic alkalies, suggested by M.O. Henry, is founded on the property of tannic acid to precipitate the organic alkalism in gausse!

Neutralize by potassa a clear infusion obtained by digesting the vegetable matter containing the alkali, or an extract produced from it, in tepid water, acidulated by sulphuric acid add an infusion of galls so long as any precipitate ensure. The precipitate, after being washed with cold water, is to be thoroughly sungled with by drate of time, somewhat in excess, and being dried by the heat of boiling water, must be digested in alcohol or other. The resulting solution, after filtration, is to be subjected to a heat sufficient to drive off the alcohol. The residual liquid, connected of water which had been in combination with the alcohol, holds the alkali in other, and after a few days repose deposits it in crystals.

and again precipitated by ammonia, may be purified of the matter by which they are, in the first instance, discoloured. A particular account of an improved process, devised by Dr. Staples, may be found in the United States Dispensatory, by the editors of which it is highly recommended.

5415. The following process, suggested by Wittstoch, is recommended as probably the best, by Kane.

5416. One part of opium, from eight to ten of water, with two of chlorohydric acid, are to be digested together for six hours. The solution being then decanted, the residue is to be subjected twice successively to the same ordeal. The resulting solutions being united, the whole is to be saturated with chloride of sodium. The matter which consequently subsides, is to be separated by filtration, and ammonia being added, in slight excess, to the filtered liquid, it must be allowed to rest undisturbed for twenty-four hours. The resulting precipitate is to be colected upon a filter, washed with a little water, dried, and digested in alcohol, of 0.820, which takes up the morphia. The greater part of the spirit being removed by distillation, morphia crystallizes on cooling in a state sufficiently pure.

5417. The effect of the chloride of sodium is to precipitate narcotina, and some other impurities. The meconin, codeia, thebaine, and some other principles, are re-

ained in solution by the alcoholic mother liquor.

5418. Morphia crystallizes in rhombic prisms, containing for each atom, two of water, which are liable to be lost by efflorescence. It has an enduring bitter taste, and is almost insoluble in water, as it requires for solution 400 parts, even at the temperature of ebullition, and precipiates, most entirely, as the liquid cools. It has an alkaine reaction, readily dissolves in alcohol, but sparingly in It is also soluble in aqueous solutions of the alkalies and earths.

5419. As usually procured, this alkali, or any of its combinations, is reddened when brought into contact with nitric acid. The phenomenon is produced by the same acid on contact with other vegeto-alkalies, and, according to Kane, is not produced with morphia when absolutely pure. Subjected to chlorine in water, morphia is first made orange red, and then dissolved. On contact with morphia, the iodine of iodic acid is liberated. A solution

of sesquichloride of iron assumes a rich blue colour on the addition of morphia, or any of its salts. With tannic acid morphia affords a copious white precipitate. It is capable of neutralizing the strongest acids, and of forming with them compounds which are soluble and crystallizable.

5420. Agreeably to the late observations of Larocque and Thibierge, the perchloride of gold produces with morphia a precipitate which is at first yellow, next bluish, and lastly violet. In the state in which it assumes the color last mentioned, the gold is revived; while the precipitate, of which it forms a part, becomes insoluble in water, alcohol, caustic alkalies, or in sulphuric, nitric, or chlorohydric acids. Yet with aqua regia, it makes a solution which is precipitated by the green sulphate of iron.

5421. With the oxacids, with organic acids, and with the halogen bodies, morphia generates salts which are capable of crystallization and of being dissolved by water. The medicinal properties of the alkali are not impaired by these combinations. In this country the sulphate is the most used; but Dr. Kane alleges the "muriate" to be the

most important compound of morphia.

# Of Paramorphia, or Thebaine.

5422. Paramorphia is an alkali lately discovered by Pelletier in minute proportion in opium. It is identical with morphia in composition, but quite distinct in its properties. It is, therefore, isomeric with morphia, and hence its name.

5423. Paramorphia is white, scarcely soluble in water, of an acrid and styptic, rather than a bitter taste, and very soluble in alcohol or other, even when cold, and still more so when hot. It differs from morphia in not being reddened by nitric acid, in not forming crystallizable salts with acids, and in not striking a blue colour with the salts of iron. It also differs from morphia in its action on the system, producing tetanic symptoms in doses of a grain.

5424. Pseudomorphia is a name given to another alkalidiscovered by the same distinguished chemist in opiumlikewise in minute proportion. It resembles morphia in the characteristic properties of becoming red with nitre acid, and of striking a blue colour with the salts of iron and yet differs from it in not being poisonous. It is not always present in opium, and the circumstances under which it is produced are not known.

# Of Codeia, or Codeine.

5425. This vegetable alkali was discovered in 1832, by tobiquet. It exists in opium as a meconate. It is in the orm of colourless crystals, which are soluble in two parts f boiling water, also soluble in alcohol and ether, but include in alkaline solutions. Its capacity of saturation very nearly the same as that of morphia; but it may be istinguished from that alkali by the different form of its rystals, by its greater solubility in water, and by its includility in alkaline solutions. It has a decided action n the animal economy, producing first excitation, and aferwards depression.

## Of Narcotina, or Narcotine.

5426. In order to obtain narcotina, opium may be comninuted, and digested with as much ether as will cover it,
t a temperature near the boiling point of the ether, for
hree or four days. The ether being decanted and allowd to evaporate, the narcotina will appear in slender prisnatic crystals, soiled by caoutchouc, resin, and colouring
natter. Being subjected to boiling alcohol and recrysallized by refrigeration therefrom, they are rendered purer,
nd further purified by repeated solution and recrystallizaion. To remove all the narcotina, opium must be subscted to successive portions of ether.

## Of Narceia, or Narceine.

5427. This alkali was discovered in opium by Pelletier n 1832. It exists in white, silky, acicular crystals, inolorous, of a slightly bitter taste, sparingly soluble in water, nore soluble in alcohol, and insoluble in ether. It is renered blue by the dilute mineral acids, but does not, like norphia, become blue with the salts of iron, nor red with itric acid.

### Of Quinia, or Quinine.

5428. In the various kinds of cinchonia, known in comnerce as Peruvian bark, there are three organic alkalies, quinia, cinchonia, and aricina, of which the most imporant is that which bears the name at the head of this paagraph. Quinia is generally procured from yellow bark. The process usually employed for its elaboration is as ollows. The bark, coarsely powdered, is boiled with sulphuric or chlorohydric acid. In the case of sulphuric acid, the proportions given are three fluid drachms to a gallon of water; in the other case, two of acid to ten of water;

pound of bark being employed.

5429. The bark is to be subsequently exposed to a similar ordeal with a half, and with a fourth part of the quastity of acid at first employed. To the united solutions, strained and cooled, add hydrate of lime till there bear alkaline reaction. The precipitate is to be collected. This, when sulphuric acid is used, will consist in part of sulphate of lime; but when the other solvent is used, the lime will remain in solution in the state of chloride. h either state, the precipitate being digested in alcohol, the alkali is taken up. The solution thus formed, is subjected to distillation with water. The residue being treated with sulphuric acid in excess, on evaporation affords crystals of sulphate of quinia; the sulphate of cinchonia remaining in solution. From the sulphate, pure quinia may be contained by adding to a solution of it caustic potash, also in solution, drying the resulting precipitate, dissolving it in a quantity of alcohol, as small as possible, and allowing the liquid thus obtained to evaporate leisurely in a place moderately warm. Under these circumstances, quina crystallizes in union with an atom of water, forming of course a crystalline hydrate. This water it loses by Quinia is intensely bitter. It requires for solution, two hundred parts of hot water, and is almost insoluble in cold water. In alcohol or ether it dissolves readily. The salts of this alkali are soluble in water, as well as m alcohol, and are capable of crystallizing. In common with those of other alkalies, and of ammonia, the oxysalts which it forms, require an atom of water, as already mentioned (5406).

5430. Of the Chlorohydrurct or Muriate of Quinia. This salt forms pearly crystalline needles, which are very soluble in water. It acts as a base with chloroplatinic, or chlorohydrargyric acid (corrosive sublimate), forming what are called double salts by some chemists, but which I corceive should be called, severally, chloroplatinate or chlorohydrargyrate of the chlorohydruret of quinia; or for the sake of brevity, as in other cases, simply a chloroplatinate of morphia, or chlorohydrargyrate of quinia (5406).

5431. Basic Sulphate of Quinia consists of two atoms

f quinine, one of sulphuric acid, and eight of water; its ormula being Qu So<sup>3</sup> 8HO. The manufacture of this ompound is conducted on a large scale, according to the rocess above given for the extrication of quinine, and vaious other methods. In crystallizing, this sulphate enters 1to combination with six atoms of water of crystallizaion, and two acting as a base. Hence in dry air, or when ently heated, it relinquishes six, yet retains two, which annot be expelled without partial decomposition. alt is but sparingly soluble in water, requiring thirty parts t a boiling heat, and seven hundred and forty in the cold. If alcohol, unaided by heat, it requires eighty parts for olution; but much less at the temperature of ebullition. ts crystals are small pearly plates or needles, which, when eated, fuse, and phosphoresce vividly, being totally decomosed at a high temperature.

5432. Neutral Sulphate of Quinia.—This salt crystalzes in rectangular prisms, of which the formula is Qu So<sup>3</sup> HO. They are prone to effloresce, dissolve in ten parts f water at 60°, and undergo aqueous fusion at 112°. This sulphate is very soluble in alcohol, and, though from

s constitution it should be neutral, reddens litmus.

tate in which basic sulphate of quinia is sold in comnerce, under the name of sulphate, it is sometimes adulteated with boric acid and with sulphate of lime. These
nbstances may be detected by exposing the aggregate to
red heat, by which the elements of the sulphate may be
issipated, and the adulterations exposed to view. Sugar
nd margaric acid have also been used as adulterations.
If these, the latter may be detected by its insolubility in
iluted acids, the former by washing a sample in water,
nd adding carbonate of soda to precipitate the quinia,
rhen the sweet taste of the sugar will become perceptible.

5434. Phosphate of Quinia crystallizes in small, but bril-

iant needles, soluble both in water and in alcohol.

5435. Ferroprussiate, or Cyanoferrite, of Quinia is formed by boiling one part sulphate of quinia, and one and a half of cyanoferrite of potassium, in seven of water. The generated salt separates as greenish yellow, oily substance. The mother liquor being decanted when cold, the cyanoferrite is to be redissolved in boiling alcohol, whence on efrigeration it crystallizes in greenish yellow needles.

### On the Reaction of Chlorine with Quinia and its Salts.

5436. If sulphate of quinia be made to form a dilute relution with water, impregnated with chlorine, and liquid ammonia be added, a green precipitate ensues, the liquid assuming an intensely green colour. The precipitated substance has been called dalleiochin. If the restoral green liquid be evaporated with access of air, it changes to dark red, while sal ammoniac is generated, and to bodies, of which only one is soluble in alcohol. The soluble body is called rusiochin, the other, melanochalkane's Elements.

#### Of Cinchonia, or Cinchonine.

micrantha) from which it may be extricated by means and logous to those employed in the case of quinia. Usually it is obtained from the mother waters of the sulphate of the alkali last mentioned, by saturating the excess of and by which it is retained in solution during the crystallization of the sulphate of quinia. Under these circumstances being precipitated by an alkaline base, and afterwards redissolved by alcohol, it is obtained in thin, colourless, prematic crystals, by vaporizing this solvent. Its taste is peculiar, as well as bitter. Boiling water only takes up part; but it readily dissolves in alcohol and ether. It fuses at 330° without loss. Between its salts and those of quinia, there is a great resemblance.

5438. The chlorohydruret of cinchonia crystallizes in brilliant interwoven needles, and like the congenerous compound of quinia (5389) acts as a base with electronegative chlorides, such as chloroplatinic, and chlorohydrargure

acid.

5439. Basic sulphate of cinchonia, C<sup>3</sup> + SO<sup>3</sup> forms rhombic prismatic crystals, which require for solution 54 parts of water. The neutral sulphate holding only half as much base, is more soluble, crystallizing in large well formed rhombic octohedrons.

#### Of Aricina, or Aricine.

5440. This alkali was discovered in 1829 by Pelicus and Coriol, in a bark brought from Arica, on the Pacific

with the Calisaya bark. It is a white, transparent, crystalline substance, having a warm and intensely bitter taste, which is long in developing itself. It dissolves in alcohol and ether, but is completely insoluble in water. By nitric acid it is coloured green. The salts agree in their properties with those of quinia and cinchonia.

# Of Strychnia, or Strychnine.

- 5441. The poisonous principle of the Strychnos nux vomica, and Strychnos ignatia or colubrina, is considered as an alkali, and called strychnia. It may be developed by a process similar to that used for morphia. It was originally obtained by Pelletier and Caventou, by subjecting the bean of the strychnos ignatia, duly rasped, to nitric ether in a Papin's digester, to remove fatty matter; and subsequent exposure of the residue to alcohol, in which the strychnia, in union with an acid, dissolves. The alcohol having been evaporated, and the residuum dissolved in water, the addition of potash caused the alkali to precipitate. It was afterwards washed in cold water, and redissolved in alcohol, from which it crystallized by evaporation.
- 5442. The colour of strychnia is white. Its taste is intolerably bitter, leaving a metallic impression in the mouth. It is nearly insoluble in water, or ether, but is very soluble in alcohol. It is a terrible poison, very small quantities producing tetanus to a fatal extent; being used by the natives of Borneo to render their arrows poisonous, under the names either of upas tieuta, or woorara.

# Of Brucia, or Brucine.

dysenterica, or false angustura. The bark was first subjected to sulphuric ether, and afterwards to alcohol. The alcohol being evaporated, afforded a dry residuum, which was dissolved in water. The solution in water was saturated with oxalic acid, and evaporated to dryness. An oxalate of brucia resulted, which, after being depurated by alcohol of colouring matter, with which it was associated and disguised, was decomposed by lime or magnesia. As either of these bases forms an insoluble salt with oxalic

acid, while brucia is soluble in 500 times its weight of boiling water, or in 850 parts of cold, it was separated

from the insoluble oxalate by water.

5444. Brucia crystallizes in oblique prisms, with parallelograms for their bases. It is less bitter than strychain, but its taste is more acrid and durable. It melts when heated a little above 212°, and congeals on cooling into a mass resembling wax. It neutralizes acids, affording a distinct class of salts. On animals, its effects are and gous to those of strychnia, but less violent.\*

# Of Delphia, or Delphine.

5445. It was in the seeds of the Delphinium staphisegria, or stavesacre, in which it exists as a malate, that the alkali was detected. A decoction of the seeds, which had been cleansed and reduced to a pulp, was filtered. The fluid, which passed the filter, was boiled with magnetic which liberated the delphia. It was then separated from the magnesia by alcohol, and from this solvent by evaporation.

5446. Delphia is white, pulverulent, and very soluble is alcohol and ether. It is inodorous, but its taste is extremely acrid and bitter. Water derives from it an acrid taste, though it does not dissolve any appreciable quantity. By combination with acids, it forms neutral salts, which are

soluble in water, and very acrid and bitter.

5447. Concentrated sulphuric acid reddens, and afterwards carbonizes delphia. Chlorine renders it green-Courbe alleges that stavesacre contains, in addition to that which has been described, a yellow, resincus substance, of which the formula is  $C^{32} H^{23} O^4 N$ ; and the name suggested for it is staphysain. This is distinguished by insolubility in ether, or water; and solubility in dilute acids, without neutralizing them.

Mr. Fuch advances that brucia is a combination of strychnia with a reservision this last mentioned substance holds obstinately, and which has the perty of being reddened by nitric acid. It is to this impurity that brucia ower liability to be made red by the acid above mentioned. Mr. Fuch has found a method of separating this resin from brucia, and consequently of converting this supposed peculiar alkali into strychnia. He has not, however, succeeded in causing stryches to combine with the resin in question so as to form brucia. Although Mr. Fish mentioned it to be his intention to publish his process for the deparation of breeze a year has clapsed without any further information having been promulgated by the on this subject. Berzelius' Report for 1841, p. 141.

# Of Veratria, or Veratrine.

5448. Veratria is an alkali obtained from the seed of ne Veratrum sabadilla; also from the roots of the Veratum album (white hellebore), and Colchicum autumnale

meadow saffron).

5449. The seeds, partially depurated by digestion with ther, yielded a coloured tincture with heated alcohol. his tincture deposited some waxy matter on cooling, nd by evaporation afforded a residuum, soluble in water, acepting a small portion of extraneous matter. The wayry solution being slowly and partially evaporated, until n orange-coloured precipitate ceased to appear, acetate flead was added to it. A copious yellow precipitate enact, and the liquor, being separated from it by a filter, ecame almost colourless. This fluid was subjected to alphydric acid, to precipitate any lead which it might ontain. The solution then gave, with magnesia, a precipate, from which alcohol took up veratria. From the alpholic solution, the veratria was afterwards isolated by raporation.

5450. Veratria is white, pulverulent, and inodorous, at, nevertheless, poisonous when inhaled, producing viont and dangerous sneezing. Its taste is not bitter, but at a cessively acrid. It reacts like an alkali, is insoluble in ater, but very soluble in alcohol and ether. It melts at 30°. Its salts are for the most part crystallizable and eutral, but decomposable by water into free acid, and a asic salt. Taken into the stomach in minute quantities, produces intolerable nausea and vomiting, and in large

oses, death.

65

# Of Sabadilla.

5451. Sabadilla was discovered by Couerbe, as an alali accompanying veratria in veratrum sabadilla, and in the roots of the Veratrum album (white hellebore), and

blchicum autumnale (meadow saffron).

5452. By boiling the precipitate, obtained by carbonate foods from an infusion of sabadilla seeds in diluted sulturic acid, sabadilla may be separated in radiated needles, fa pale rose colour, which may be rendered white by detaition. This alkali is a white, crystallizable substance,

insupportably acrid, fusible by heat, readily soluble in be water, very soluble in alcohol, and wholly insoluble ether.

#### Of Jervina, or Jervine.

with veratrine, from which the sparing solubility of sulphate, and its readiness to crystallize from an alcoholution with four atoms of water, renders it liable to separated. Jervina, when pure, is white, easily fusible, composable at 400°, nearly insoluble in water, but copiedly soluble in alcohol. Of its salts, the acetate readily solves in water, although in this liquid its sulphate, not and chloride, are sparingly soluble. The chloride of principles with chloroplatinic acid. Kane, 1069.

#### Of Colchicina, or Colchicine.

5454. Colchicina is a vegeto-alkali existing in the

of the meadow saffron (Colchicum autumnale).

Digest the seeds in a mixture of sulphuric acid and was alcohol; neutralize the excess of acid by lime, remove alcohol by distillation, decompose the residual liquor by a bonate of potash in excess, dissolve the washed and do precipitate in absolute alcohol, decolorize the solution animal charcoal, add a few drops of water, and evaporate that the colchicina crystallizes in colourless needles.

the taste as veratrine, nor is it productive of violent such ing. It is moderately soluble in water, very soluble in a cohol, or ether. Though but feebly alkaline in its reaction in other respects it neutralizes acids thoroughly. By time ture of iodine it is precipitated of a rich orange colour, a nitric acid it is coloured dark violet blue. Though not abundant in the seeds, it pervades all parts of colchism Kane, 1069.

#### Of Emetia, or Emetine.

5457. This alkali is obtained from ipecacuanha. The roots, well pulverized, are digested in other. They then subjected to alcohol, the resulting solution is evaporated, and the residuum dissolved in water, and macerated upon magnesia, which causes the emetia to precipitate

This precipitate is washed with cold water to remove colouring matter, and afterwards subjected to alcohol, which takes up the emetia. The emetine again separated from its solvent by evaporation, being dissolved by diluted acid, and blanched by animal charcoal, may be precipitated pure

by any of the alkaline oxides.

5458. Thus obtained, emetia is white, pulverulent, and unalterable by the air, scarcely soluble in water, but very soluble in ether or alcohol. Its taste is slightly bitter. It possesses strong alkaline properties, restoring the colour of litmus, when reddened by an acid. It is capable of forming salts, which, though neutral, are not crystallizable. It appears to possess all the emetic properties of the root from which it is procured.

## Of Solania, or Solanine.

5459. Solania is the name which has been given to an alkali which exists in the black nightshade (solanum nigrum), and in the bittersweet (solanum dulcamara), also

in the shoots of the solanum tuberosum, or potato.

5460. The filtrated juice of the berries of the nightshade being digested in ammonia, the resulting precipitate is washed on the filter, and digested in boiling alcohol. After the evaporation of this fluid, solania is obtained in sufficient purity. It is a white, opake, pearly powder, which is inodorous, slightly bitter, and nauseous. Its acid solutions are more bitter. Its salts, though neutral, are uncrystallizable. In cold water it is insoluble, and in hot dissolves only to a small extent. It is very soluble in alcohol, but is not dissolved by ether. It restores the colour of litmus, reddened by an acid. It causes vomiting at first, afterwards sleep, or death, according to the dose, being a strong narcotic poison. With salts of emetine, tannic acid, or corrosive sublimate, it produces white precipitates; with iodine and chloroplatinic acid, brownish yellow preci-According to Kane, the injurious properties of unripe potatoes result from the presence of this body. It exists abundantly in the early shoots (underground) and buds of the tubers.

# Of Caffein or Caffeia,\* or Theine.

5461. It seems hardly credible that there should be a crystallized nitrogenated principle common both to team to coffee. Yet, agreeably to analyses recently made, to substances which had been discovered in tea and coffee, and called theine, or caffein, are identical in composition and properties.

5462. Moreover, a principle elaborated from guarde, a paste made from the seeds of paullinia sorbilis, is alleged by Martius to be identical in composition with caffein,

to be a base in its properties.

5463. To extract caffeia, the raw coffee seeds, well dried and pulverized, are to be exhausted by boiling water. In the next place subacetate of lead must be added to the resulting solution. This is to be filtered afterwards, and any excess of lead precipitated by sulphydric acid. After a second filtration, the solution being concentrated sufficiently by evaporation, the caffeia crystallizes on cooling-Re-solution, and recrystallization are requisite to render it pure.

5464. Caffeia may also be extricated from a filtered de-

coction of tea leaves: hence its other name, theine.

dles, having a silky lustre. It is feebly bitter, sparingly soluble in ether, cold water, or alcohol. At 212° it loses eight per cent. of water. It fuses at 352°, and sublimes at 725°. From its solution it may be thrown down by tannic acid. Boiled with caustic potash, or baryta, caffeia is resolved into ammonia, cyanuric, formic, and carbonic acids. With sulphuric or chlorohydric acid it forms crystaline compounds. Its composition, according to Liebig, is represented by the formula above given.

5466. Graham alleges that the active properties of mand coffee are not due to caffeia; but it is admitted that no other vegetable substance contains so large a propor-

or to classify, on account of the discordancy of the authorities which bear upon to classify, on account of the discordancy of the authorities which bear upon to question. Heretofore it has been placed among the neutral principles, and in the United States Dispensatory, and in the recent works of Kane. Graham and Grapery, has been treated of as such, and called caffeia. But in the report of Berreis for 1841, it is mentioned that Martius has a found it to be identical with guesan rean organic base, elaborated from the seeds of paullinia sorbilis. Accordingly it placed by Berzelius, in his list of contents, under the head of vegetable tasks at morphine, brucia, &c. But while Martius and Berzelius assign to it the rank of a alkali, they do not change the terminating monosyllable, as the continental change have not adopted the termination in a for alkaline bases.

tion of nitrogen, and Liebig remarks that  $2\frac{1}{10}$  grains of caffeia may turnish all the nitrogen required by an ounce of human bile. This fact naturally suggests that tea and coffee may be serviceable in furnishing nitrogen for biliary and other secretions, in beings whose habits of life do not make it healthful or agreeable to consume a sufficient quantity of bread and meat to supply all the nitrogen necessary to the vital functions.

markable, that civilized nations, comprising a majority of markable, that civilized nations, comprising a majority of markind, should in modern times have been led, as it would seem, intuitively, to resort to two sources, apparently so different, as the tea leaf and coffee berry, for the same preëminently nitrogenated principle as an almost indispen-

sable article of daily food.

# "Chelerythrina, or Chelerythrine.

5468. "This substance is extracted from the roots of the chelidonium majus, by digestion with dilute sulphuric acid. The liquor so obtained is to be evaporated and mixed with ammonia. The brown precipitate which falls is to be washed, pressed between folds of paper, and digested in alcohol, with some sulphuric acid. The alcoholic solution being mixed with water, and the spirit distilled off, the residual liquor is precipitated by ammonia, and the precipitate being washed and dried by pressure, is to be digested in ether, and the ethereal solution evaporated to dryness. The mass so obtained is then digested in dilute muriatic acid, which leaves a resinous substance undissolved. The deep red liquor evaporated to dryness, and washed with ether, leaves a mixture of muriate of chelerythrine and muriate of cheledonine; the former of which is dissolved by washing with a small quantity of water, whilst the latter remains undissolved."

5469. "From the solution of the muriate, the chelerythrine is precipitated by ammonia, as a white curdy powder. From its ethereal solution it remains as a resinous mass, which remains soft for a long time; it is insoluble in water; its solutions in alcohol and ether are pale yellow. With acids it forms salts of a rich crimson colour, which generally crystallize. Tannic acid produces in their solutions a precipitate soluble in alcohol." Verbatim from Kane, 1070.

#### "Chelidonia, or Chelidonine.

great part described in the preceding article. By desing the sparingly soluble muriate with ammonia, the solving in sulphuric acid and precipitating with main acid, it is freed from all traces of chelerythrine, and the pure chelidonine, separated by ammonia, is distributed in boiling alcohol, from which it crystallizes, on continuous in brilliant colourless tables. It is insoluble in water, buble in alcohol and ether; it tastes bitter, and real acids crystallize; its solutions give with the main a precipitate." Verbatim from Kane, 1071.

#### Of Atropia, or Atropine.

leaves of the Atropa belladonna, or deadly nights. Two pounds of the leaves were boiled in successive tions of water, which being united, and sulphure added to the whole, the resulting liquid was filtered, yielded a crystalline precipitate with potash. This printed, repeatedly dissolved in acids, and precipitate alkalies, gave pure atropia. Thus obtained, it is white, and quite tasteless. When recently precipitate is slightly soluble in water. After being dried, it is not ble in water, ether, or oil of turpentine. In cold alcohol is sparingly soluble; but copiously in the same mension when boiling hot.

5472. Atropia forms compounds with acids, which a not, however, be rendered so neutral, as not to indicacidity.

#### Of Aconitia, or Aconitine.

5473. The fresh expressed juice of the monkhood, a nitum napellus, being boiled and filtered, the resulting a liquor, subjected to an excess of carbonate of potash, it be agitated with ether so long as it takes up any this On vaporizing the ether, aconitia is deposited. From dry plant, or its seeds, a solution of aconitia may be tained by water holding an ounce of sulphuric acid each pound. This may be decomposed by carbonate

nd the alkali extricated from the resulting precipiy ether or alcohol. Aconitia crystallizes from an l or alcoholic solution, partly in white grains, but most part forms a colourless vitreous-looking mass. a sharp bitter taste, and is intensely poisonous. It ble of neutralizing the most powerful acids. Its sogive a white precipitate with alkalies proper, or aloride of gold; with iodine an orange precipitate.

# Of Belladonia, or Belladonine.

Let This alkali is obtained by subjecting the dried belladonna to distillation with a solution of caustic, precipitating, from the liquid which comes over, ali with which it is accompanied, by chloroplatinic and heating the washed precipitate with carbonate ash. The belladonia being sublimed, condenses in ess, rectangular, prismatic crystals. Belladonia, olated, has a penetrating odour resembling that of ita, and forms a solution with water, which reacts at of an alkali. It is not very poisonous. Its salts ich like the corresponding ammoniacal salts.

# Of Daturia, or Daturine.

i. The seeds of the datura stramonium, vulgarly as the thorn apple, Jamestown, or jimson weed, e juice of the leaves, capsules, and stems, contain taline principle to which the name at the head of ticle is given. It is to this, that the efficacy of the nt constituted by the inspissated juice, and the well poisonous property of the plant, are due.

Agreeably to the process of Brandes, who first daturia, the seeds are to be boiled in alcohol, and sia being added, the resulting precipitate is to be lived by the same liquid. According to Kane, it obtained by the same processes as aconitia, above sed.

7. From its solution in spirit, it crystallizes in very it groups of needles. It is quite inodorous when although the juice of the plant smells disgustingly ic. It is bitter, and tastes somewhat like tobacco. solution, it requires 72 parts of boiling water, 250 water, 21 parts of ether, and 3 of alcohol. It fuses 212°, and at a higher temperature volatilizes, un-

changed, in white clouds. It reacts like an alkali, and capable of forming, with acids, crystallizable saits, where highly poisonous. In its habitudes with reagent resembles atropia.

#### Of Conina, or Concine.

5478. This alkalı exists in all parts of the hemlocking nium maculatum), especially in the seeds, from which may be extricated by the following means:—They are be bruised, and being mingled with one part of a conc trated solution of potash, and eight of water, are to be jected to the distillatory process till the water, which calls, becomes inodorous. The distilled solution, after be neutralized by sulphuric acid, must be evaporated to 1 consistency of a syrup; and being, in this state, tres two or three times with a mixture of one part of ether, two of alcohol of 820°, the coneine is taken up. water being added, the ether and alcohol are removed distillation, and the residual water by evaporation. The siccated residuum is to be mingled with half its weight a concentrated solution of caustic potash, and subjected distillation with a receiver carefully refrigerated. portion must be separated from the aqueous portion of liquid which comes over, and this last again distilled 🛍 hydrate of lime. From any ammonia with which it be associated, the coneine may be freed by exposure for few hours in vacuo, over sulphuric acid.

5479. Pure coneine is extremely poisonous, existing the form of a colourless transparent liquid, of the density ( .890. Its taste is disgustingly sharp, its smell highly and seous and pungent, somewhat like that of the plant. It is soluble in 100 parts of cold water, which becomes turbi by being heated; but four parts of coneine dissolve oned water, forming a solution which may be rendered tubb by the heat of the hand. With alcohol, ether, and oil, i mingles in all proportions. It distils, per se, at 370°, in requires less heat when associated with the steam of bo ing water. It reacts like an alkali with the assistance t water, but not when anhydrous. It is capable of set rating acids completely, having the least atomic weight any known organic alkali. Its salts, which crystallize in imperfectly, are decomposed by much water. In alcoho or a mixture of this solvent with ether, they readily di olve, but are insoluble in pure ether. The precipitate iven by their aqueous solutions with iodine is saffron yel-

w; that yielded with tannic acid, white.

5480. Coneia is coloured blood-red by nitric acid. By rposure to air it turns brown, and is resolved into ammoia, and a bitter, inodorous, resinous substance, which is poisonous.

# Of Nicotina or Nicotine.

5481. The preceding name is given to the active poinous principle, to which tobacco (nicotiana tobaccum) and some other plants owe their active qualities. For its aboration, the means described as suitable for the elabotion of coneia may be used, though in either case magnia, or any other alkaline earth, or alkali, might be sub-

ituted for potash in the first step of the process.

5482. Pure nicotina or nicotine is a colourless oily liquid, dowed, in a high degree, with the odour and taste of toicco. It is soluble in water in all proportions, which is
property displayed by no other organic base. It is also
luble in ether or alcohol. When anhydrous, it emits
hite fumes at 212°, and at 480° distils, undergoing, howrer, a partial decomposition. Its distillation is accomished easily with the aid of water.

5483. Nicotina is highly alkaline, neutralizing and formg soluble salts with acids. Of these, some are crystallible, retaining, however, the savour of tobacco. Subcted to alkalies, they evolve the characteristic odour of

e plant.\*

5484. Of Lobelina or Lobeline.—It appears by an article the American Journal of Pharmacy for April, 1841, Vol. 1, that Mr. Procter, jr., has obtained an organic alkali om the seeds of the lobelia inflata, by acidulated alcohol, splacement, ether, and evaporation. This alkali is repre-

A new process for the evolution of nicotina is given in the Journale de Pharma
1, for February, 1842, of which the steps are as follows:—Maceration for 24 hours

water acidulated by sulphuric acid; expression, evaporation to a syrupy consist
ce; distillation with potash, water being added to prevent injurious concentration;

straigntion by oxalic acid; evaporation to dryness; treatment with absolute alco
l, which takes up oxalate of nicotina; evaporation, decomposition by potash; solu
a in other; evaporation, whence results nicotina free from all impurity, excepting

the and alcohol in a minute proportion. Agreeably to M. V. Ortigosa, the author

this new process, nicotina forms compounds with chloroplatinic, and chlorohydrar
the acid.

sented as having a great resemblance to nicotina, but much less poisonous.

#### Picrotoxine or Picrotoxia.

dicus has received the name of picrotoxine, but he at been conceived to have basic properties, nor to agree the organic alkalies in holding nitrogen as an cleant. Nevertheless, in the late work of Liebig and Grant 1168, it is arranged among the organic bases, and be leged to have been shown, by the recent researches of Francis, to contain 1.38 per cent. of nitrogen. Yet a not formula for picrotoxine had not been published by the chemist.

5486. I subjoin an account of the process for obtaining

this alkali, and a description of its properties.

jected to pressure in order to expel as much as possible their fat oil, are boiled in alcohol. The alcohol being parated from the matter which it takes up by distillated this matter is redissolved in boiling water, slightly and lated. From the resulting solution, on cooling, the protoxine separates in short, thin, colourless prisms, insured tible of fusion. Picrotoxine is soluble in twenty-five part of boiling water, and very soluble in alcohol. It is tensely bitter, and highly poisonous. Its formula is probably C<sup>13</sup> H<sup>7</sup> O<sup>3</sup>N.

#### Of Antiarine or Antiaria.

has been given, is in a predicament analogous to that a which picrotoxine has heretofore been placed. I must that of resembling many of the organic bases in its activity as a poison, while devoid of nitrogen, and of the bility to react like a base. It is not, however, improbable that further researches may prove the pretensions of the triarine to rank with the organic alkalies, both as to prepare and composition. Antiarine is the active principle of that most deadly upas poison, respecting which, high exaggerated accounts were published about forty vestago, representing that the tree producing it could be without loss of life, be approached, unless upon the wind

urd side. Its formula is alleged to be C<sup>12</sup> H<sup>10</sup> O<sup>5</sup>. It restallizes in small scaly crystals, soluble in 250 parts of ld water, 70 of alcohol, and 2790 of ether.

# Bases from the Oil of Mustard.

5489. Thiosinnamina.—When the oil of mustard is nught in contact with three or four times its volume of ong ammonia, crystals are formed, which are purified recrystallization. These are thiosinnamina: formula, H<sup>8</sup> N<sup>2</sup> S<sup>2</sup>.

5490. Thiosinnamina is soluble in hot water, less so in ld water, soluble in alcohol and ether. It has a bitter ste, and no smell. At 392° it is resolved into ammonia, d a resinoid basic compound not fully investigated inosinnamina combines with acids, but its salts do not ystallize: it yields a chloroplatinate with chloroplatinic id = C° H° N° S° HC Cl + Pt Cl; and with corrosive subsate a chlorohydrargyrate = C′ H′ NS Cl + Hg Cl.

5491. Sinnamina.—This new base is obtained in the folving way:—Thiosinnamina is digested with moist hyated protoxide of lead till all the sulphur is removed. ne residue is then subjected to water, finally to alcoholne resulting solution is evaporated to a syrup, which, er some time, deposits fine transparent crystals of sinmina.

5492. Sinnamina is a powerful base, expelling ammonia m its salts, and precipitating the solutions of peroxide of n, of copper, and of lead. It combines with acids, but elds no crystallizable salts. It is precipitated by chlorotinic and chlorohydrargyric acid, and throws down silt from its solution in nitric acid. When heated, it pleas ammonia, and leaves a basic resinoid matter production of sinnamina from thiosinnamina is efted by the separation of all the sulphur with more or hydrogen. I say more or less, since it is not known the certainty whether the formula of sinnamina is C<sup>8</sup> H<sup>6</sup>N<sup>2</sup>. (Varrentrapp and Will.)

5493. Sinapolina.—This compound, discovered by Sin, is obtained by depriving oil of mustard of its sulpr, by the action of baryta or of oxide of lead. It is uble in hot water and alcohol, and crystallizes in ning, fatty, fusible scales. Its solution has an alkaline ction. It combines with acids, and may be separated

from them by ammonia. When combined with chloroly-dric acid, it precipitates the chloroplatinic and chloroly-drargyric acids. It is generated from the oil of mustard by the abstraction of two atoms of bisulphuret of carbon and the addition of two atoms of water. Thus C<sup>16</sup> H<sup>18</sup> N S<sup>1</sup> + 2HO = C<sup>14</sup> H<sup>18</sup> N<sup>2</sup> O<sup>2</sup> + 2CS<sup>2</sup>. The formula of single-polina is C<sup>14</sup> H<sup>18</sup> N<sup>2</sup> O<sup>2</sup>. Liebig and Gregory, 1156.

because of the alleged inefficacy of the species of cinchona, from which is derived, it may be inferred that cinchovine has little or no practical value and will not merit that more should be said of it here. Comptes Read 25, 125.

5495. Of Cisampelina or Cisampeline, also called polosine. In his its port on Chemistry for 1841, Berzelius gives the following information as specting this base, lately discovered by Wiggers. A filtered solution of the roots of cisampelos pareira, obtained by digestion in water acidalated in sulphuric acid, is saturated with carbonate of soda, avoiding to add as a cess. The precipitate twice washed, and well dried by filtering paper. It subsequent exposure to a heat of 212°, is subjected to pure ether. It is recovered from it pure and anhydrous by the distillatory process.

5496. Cisampelina, thus procured, is hard and brittle, and to the massweetish bitter and nauseous. It has not been crystallized. It is to the principle that the medicinal properties of the cisampelos pareira are cribed. The alkali is called pelosine by Wiggers, its discoverer; but I concur with Berzelius, that the other appellation is preferable as reading the idea of its source.

tenschmidt alleged that he had discovered two bases in the "cortex guarm jamaicensis and surinamensis." Agreeably to Berzelius' report, the extense of these bases has lately been confirmed by Wiggers. Vandames and Chevalier, according to the same authority, have discovered a base in hedera helix. As no important efficacy is ascribed to these bases, I do not deem it necessary to notice them further. The same considerations have prevented me from noticing some other bases, of which accounts are the found in the reports of the great Swedish chemist; and likewise industrial ammeline, derived from melam, a product of the decomposition of subphocyanide of potassium.

Of certain general characteristics of the Vegetable Alkalies distinguishing them from Inorganic Bases, and of those which distinguish the into several different sets.

5499. It is observed by Liebig and Gregory, that the organic bear of quire less acid for saturation in proportion as they contain more oxygenalthough it is well known, that the more the oxygen in an inorganic has the greater the quantity of acid which its saturation requires.

5499. Agreeably to the same authority, the salts formed with aconita, atropia, brucia, cinchonia, codeia, conicina, delphinina, emetia, morphia, parcotia, quinia, strychnia, veratria, are precipitated white by an infusion of galls. The precipitate is a tannate, which, by exposure to the air, becomes converted into a soluble gallate.\*

5500. I will here quote from Liebig and Gregory the following arrangement of the alkalies, as I consider such generalization always instructive,

and serviceable to the memory.

1. Volatile bases containing no oxygen.

These are anilina and nicotina, to which may be added conicina, although it is not certain that this base is destitute of oxygen.

2. Bases derived from the oil of mustard.

These are thiosinnamina, sinnamina, and sinapolina.

3. Bases of cinchona bark.

These are quinia, cinchonia, and aricina.

4. Bases of the papaveraceæ, or the various species of poppy.

These are morphia, codeia, narcotina, thebaina, pseudomorphia, narceia, and chelidonia.

5. Bases found in the solanaceæ, strychnaceæ, and other plants of the same kind.

These are atropia, solania, jervina, brucia, strychnia, sabadillia, veratria, delphia, staphisia, menispermia, picrotoxia, emetia, corydalina, berbina, pi-

perina, harmalina, caffeia, and theobromia.

when chlorohydric acid combines with an organic alkali, it can form a combination meriting to be called a chlorohydrate, while the compound which is engendered by the contact of this acid with ammonia is supposed to be ammonium; in other words, a chloride of the hydruret of that gaseous body. On the subject of iodine, Berzelius has urged that "a direct combination of it with a vegetable alkali is as unlikely to exist, as would be a like combination with any other salifiable base; and, moreover, experience hows, that such compounds are neither iodates, nor iodohydrates of the vegetable alkali."

5502. It must be evident, that whatever objections exist to assuming the existence of iodohydrates of organic bases, apply with equal force to the

existence of chlorohydrates, bromohydrates, fluohydrates, &c. &c.

measure, on the investigations of Bouchardat, the vegetable alkalies have, in common with ammonia, a propensity to combine with two atoms of ioline, the recognised combinations consisting, not of an atom of iodine and in atom of the vegetable alkali, but of a compound of iodine and an iodorydrate of such an alkali. This view of the subject is alleged to be coroborated by the fact, that the combinations, with organic bases alluded to, are obtained, with pre-eminent facility, by a double decomposition consequent to the reaction of bi-iodide of potassium with a salt formed by an acid with one of these alkalies. The precipitates of the alkalies in question, thus altained, are nearly insoluble, and in many instances well characterized. Ience the bi-iodide of potassium may be more confidently relied upon as a recipitant of the organic bases than tannic acid. From the precipitated

This does not altogether confirm the allegation quoted from O. Henry (5307, etc.), that tannic acid may be used as a general mean of precipitating, and thus obtaining the vegetable alkalies. No suggestion is made as to any advantageous meads of extracting the alkali from the precipitate.

compound of iodine with the organic base, the latter may be liberated by subjecting them in water to sulphydric acid. By these means the sodes a converted into iodohydric acid, after which, an inorganic alkaline base of separate the organic alkali in an isolated state. Berzelius' Report for 1949, p. 179.

5504. It may be proper to mention, that bi-todide of potassium is formal by digesting rading in a solution of iodide of potassium, usually expressory designated in the shops as hydriodate of potash. The bi-rodide cas only

exist in solution, according to Berzelius.

## Constitution of the Organic Alkalies.

5505. All the organic alkalies are constituted of hydrogen, carbon, oxygen, and nitrogen, except melamine, nice tina, and anilina, which are devoid of oxygen.

5506. It is remarkable that these alkahes contain a very large proportion of carbon, and that in all of them nitro-

"Chloride of Gold as a test of certain Vegetable Alkalies — MM Larocque tell Thibierge find, that perchloride of gold is a more decisive test of cortain regulable alkalies, than the double chloride of sodium and gold already employed for the property of the precipitates which it produces and the salts of the annexed alkalies dissolved in water - Quinia, buff coloured , conclusion, sulphur-yellow; stryclinia, conary yellow, veratria, slightly greenish-rether to, milk collection and then concentrations, morphia yellow, then blush and may violet. In this last state the gold being reduced the precipitate is cased the in outer afcohol, the caustic alkalies, and sulphiere nitric, or hydrochloric acide, but famili with aqua regia a solution which is precipitated by protosulphate of iron

"All these precipitates with the exception mentioned, are very soluble in alectal, insoluble in ether, and slightly soluble in water. They appear to be combinated gold, chlorine, and the vegetable alkali, since their alcoholic solutions, treated will tannin, give a greenish blue precipitate of reduced gold if the solution is the and the alcohol be avaporated by heat, a precipitate of tannate of the athan control in formed. The liquor again filtered, gives with nitrate of silver a white programs

insoluble in nitric acid, but soluble in ammonia. "Among the reactions of chloride of gold, those which occur with morphs and brucia, to the authors appear to be especially important, as they are sufficiently marked to prevent these alkalies from being mustaken for each other, and also yall pretty good characteristics for distinguishing brucia from stryclimia. "The authors have also, as the results of their experiments, arrived at the follow

ing conclusions — if tat. By the aid of reagents it is possible to determine the presence of marks. atrychnia, and brucia, in substances, which, after being mixed with the mits of the alkalies, have undergone the vinous, aceta or patrifictive fermentation. At the has already shown that the patrefactive fermentation does not after morphia

"2dly Crystallized indic ucid, or a concentrated solution of this acid is same ble of being decomposed by neutral azotized bodies, but a dilute solution of the cannot be decomposed by them, unless there be added concentrated susphers and orystallizable acetic acid, oxalie, citric, or tartaric acid

Gdly. Iodic acid should not be employed as a test of morphia without the present

enution

"4thly. Perchloride of gold produces such effects with the vegetable alkalisa. serve to distinguish in orphia, brucia, and strychnia, from each other

"hthly The reagents, on which the greatest reliance may be placed as torse morphia, are nitric acid neutral perchloride of iron, and perchloride of gold "othly. By the use of reagents, morphia, which has been mixed with beer,

or milk, may be detected "7thly. It is also easy to prove, by reagents, the presence of mecome acid is --or milk, especially when the meconate of lead is decomposed by dilute sulphine acid." Journal de Chimie Médicale, Octobre, 1842 (5265).

gen is likewise a constituent. It was at one time alleged, that agreeably to the analysis of Liebig, in an equivalent of any of the alkalies of this class, only one atom of nitrogen could be found; but subsequent observation has shown that this rule has exceptions, since strychnia and brucia are found each to contain two atoms of the element in question; and in some other organic bases, the proportion of nitrogen exceeds that of an atom to each equivalent.

5507. As morphia differs from codeia only in having one atom more of oxygen; and as the three alkalies of Peruvian bark differ only in the same way; quinia having one atom of oxygen more than cinchonia, and aricina one atom more than quinia, the idea has been suggested, that in either case a compound radical may exist, capable of different degrees of oxidation: hence morphia might be a bioxide, and codeia a protoxide, of the same radical; and in like manner cinchonia might be a protoxide, quinia a bioxide, and aricina a trioxide, of one radical. But were such the case when presented to chlorohydric acid, these oxides should severally have their basic oxygen replaced by as many atoms of chlorine, which is alleged not to arrive when the experiment is tried. They all form muriates, so called, under the circumstances alluded to, or chlorohydrurets, agreeably to the view which I have taken respecting their composition (5406). See Kane, 1078.

#### OF IMPORTANT NEUTRAL ORGANIC PRINCIPLES.

Of Salicin, a neutral Principle, and of some Compounds derived from it, or to the production of which it contributes.

5508. The discovery of an analogy, if not an identity, between the properties of the oil of gaultheria, and that of spirea ulmaria, induces the idea that there may be essential oils in other vegetables of the United States, which may be worthy of examination. Under these circumstances, every fact connected with the origin of the oil of spirea ulmaria, must be interesting to the lover of science. I have, therefore, deemed it expedient to give some details especting salicin, the principle from which the artificial hydruret of salycyl," saliculous acid, is extricated, and kewise of some substances resulting from the reaction of salicine with other bodies (5321, &c.).

5509. Salicin, C42 H23 O16 + 6HO. This interesting principle, discovered by Le Roux and Buckner, is found in the bark and leaves of bitter willows, and in that of some species of poplar. It is obtained by subjecting the bark, in a divided state, to successive portions of boling water. The resulting decoctions being united and concentrated by further ebullition, are, while boiling, mingled with litharge gradually added until the liquor becomes colourless. The lead, combining with the salicin, may be precipitated from it, together with various purities, by adding sulphuric acid at first, and then set phide of barium. With the aid of charcoal, and repeated crystallization, the salicin is obtained finally in delicate, silky white transparent needles, permanent in the air. It is bitter and inodorous, but without any reaction with vegetable colours. It sustains no loss of weight at a boiling heat, but at a higher temperature is decomposed, becoming yellow, resinous, evolving inflammable vapour, and finally leaving a carbonaceous residue. It is soluble in five perts of cool water, and in any proportion in boiling water. It is no less soluble in alcohol, but is insoluble in ether, or the fixed oils. It forms with concentrated sulphuric acid a blood-red solution, which is blackened when heated Any bark which contains salicin is liable to be reddened by contact with sulphuric acid. Salicin is thrown down from any of its solutions by acetate of ammonia. saliculous acid is evolved by distilling salicin with sulphuric acid and bichromate of potash, has already been mentioned (3066, 5320).

5510. Saliretine, C<sup>30</sup> H<sup>15</sup> O<sup>7</sup> + HO, is a resinous substance produced by boiling salicin either in diluted sulphuric, or chlorohydric, acid. It is solide in caustic alkalies, excepting ammonia; likewise in alcohol or other, but is insoluble in water. By sulphuric acid it is changed to a blood-red; and a seems likely that it is to the generation of this resin that the reddening of salicin by that acid is due. One atom of hydrated saliretine, with an atom of raisin sugar, comprise the elements of one atom of hydrated salicin.

5511. Chlorosalicine, C42H25Cl4O22. When a solution of salicine is impregnated with chlorine, a crystalline deposition ensues, which dissolves a water with difficulty, but in hot alcohol with ease. It may be considered comprising the same elements as salicin, excepting the substitution of the

atoms of chlorine for a like number of hydrogen.

5512. When during the impregnation, in the process above described the temperature is raised to 140°, a compound is obtained in which severations of hydrogen have been replaced by a like number of chlorine; are mula C48 H18 Cl7 O18.

5513. Rutiline. Under this appellation Braconnot designates a substance rising from the decomposition of salicine by concentrated sulphuric acid. Ture rutiline, when moist, appears at first reddish-brown, but soon becomes rellow; when desiccated, its colour is brownish-black. It is friable, inside, inodorous, and insoluble in water or alcohol. By inorganic acids its

sue is changed to a beautiful red, by alkalies to a deep violet.

5314. Phloridzine, Cas Has Ota + 6HO. The preceding name has been given to a principle discovered by De Koninck in the bark of the roots of sple, pear, cherry, and plum trees. In composition and properties it is ery analogous to salicin; and differs, as respects elementary constituents, ally in having two more atoms of oxygen. Phloridzine is extracted from my bark in which it may exist, by boiling alcohol of the specific gravity of B50. From the alcoholic solution thus obtained, it crystallizes on the renoval of the solvent in delicate, colourless, silky, rectangular, prismatic medies; which are soluble in 1000 parts of cold water, and in every proportion in boiling water. The solution has an astringent, bitter savour, without any power to change vegetable colours. In alcohol it is also soluble, but is insoluble in ether. At 212° it loses four atoms of water of crysallization. It melts at 320°, but is not decomposed under 390°.

5515. Phloridzeine, C48 H29 O26 N2. This name is employed to designate a substance obtained by the reaction of phloridzine with animonia and tmospheric oxygen. As its name differs from that of this last mentioned abstance only in the presence of an additional e, and conveys no idea of s composition, it seems very ill chosen. By simultaneous contact with atnospheric oxygen and gaseous ammonia, moist phloridzine is transformed nto a red matter, which, readily dissolving in liquid ammonia, may be preipitated therefrom by acids. The precipitate, thus obtained, is phlorideine. It is formed by the addition of eight atoms of oxygen, and the elenents of two atoms of ammonia, to phloridzine. An ammoniacal solution f phloridzeine, evaporated within an exhausted receiver, including some agments of the hydrate of potassa, is converted into a purple blue resi-uum, having a cupreous metallic brilliancy. This residuum is unalterable a dry air, soluble in cold water, to which it communicates a magnificent urple blue. This solution is decolorized by deoxidizing substances, but esumes the oxygen thus lost, and the blue colour, on being re-exposed to he air. This blue residuum is compounded of an atom of phloridzeine, nd an atom of ammonia.

5516. Asparagine, asparamide, altheine, agedoile.—These are the syno-ymous appellations of a principle capable of forming a crystalline hydrate, if Ho Oo No + 2HO, which loses its water of crystallization at 248°. It found in asparagus, in liquorice, in the root of althea officinalis, in that of se potato, and various other plants. It crystallizes in large, transparent, ight rhombic prisms. It has a cooling and somewhat nauseous taste, is oluble in water and diluted alcohol, but insoluble in this last mentioned iquid when concentrated, or in ether. By reaction with acids or alkalies, assisted by heat, asparagine is resolved into ammonia, and an acid called spartic. The considerations which were mentioned as giving importance of caffein, must apply to asparagine as being a highly nitrogenated principle, since such principles, without any very sensible activity, may, agree-bly to the suggestions of Liebig, be of importance in supplying the nitroten requisite to facilitate the functions of life.

5517. Taraxacine.—Mons. Polex has extracted from the milky juice of leontodon taraxacum, a crystallizable substance, which he has named

which means the albumen is coagulated, involving the resul, fatty many and caoutchouc. The concentrated liquor is filtered, and allowed in rate spontaneously in a place moderately warm. The taraxacus cristallizations from alcohol or water. It forms arborescent or street, and have a bitter admin acrid taste. They are sparingly soluble in cold water, but described acrid taste. They are sparingly soluble in cold water, but described antily in boiling water, in alcohol, or other. They describe in the centrated acids without being decomposed. Taraxacuse contains the gen.

the water, is boiled in alcohol, a colourless substance, in the form of make cauliflower crystals, is obtained on the evaporation of the alcohol. It is insoluble in water, but very soluble in alcohol and other. The make tion has an acid taste, and yields no precipitate with acetate of less in insoluble in the caustic alkalies. Berzelius' Report on the Proposition

Science.

#### Of certain Vegetable Principles devoid of Nitrogen.

5519. I have quoted verbatim, from Gregory and Liebig's new classical Turner's Chemistry, 1118, the following account of vegetable practical particles as devoid of nitrogen, and of a nature not yet fully ascertain in hopes that some of my pupils may be induced, by their investigation endeavour to remedy the imperfection in chemical science thus admirable exist.

butes, and purified by solution in alcohol. It forms golden velocerates of a very buter taste, which may be sublimed. According to Transmitted when quite pure it is no longer buter, and has acid properties, expensive bonic acid from the alkaline carbonates, and forming, with the distance.

golden-yellow crystallizable salts.

5521. "Santonine is found in the flowers of several specks of him miria, and in the so called Semen Cyna, which is much used us to miluge, and is a mixture of the flowers, buds, and unrape scaled " plants. Four parts of this mixture are digested with one-half of said lime and twenty of alcohol, at 90 per cent. The santonine is district a combination with lime and with a brown resin. It is separated by some acid, but is still contaminated with resin. This is removed by with a little alcohol; and the residue being dissolved in eight or ten period alcohol at eighty per cent., and boiled with animal charcoal, the or ... cooling, deposits santonine in colourless crystals, which must be ket a less dark, as they become yellow when exposed to light. It is tasteles in a odorous, fusible and volatilizable, sparingly soluble in water, more cont alcohol and ether. It has acid properties, and forms salts with potash and soda, the latter of which crystallizes. Acids dissolve it without alternat and water precipitates from the solution the santonine unchanged. It form crystalline salts with time and baryta, and inscluble compounds with mill metallic oxides. Its composition is represented by the formula C Hell (Ettling); but its atomic weight must be twelve times greater, to judge time its capacity of saturation.

22. "Picrolichenine.—Discovered by Alms in the lichen Variolaria a, from which it is extracted by alcohol. It is purified from a green which accompanies it, by washing with a dilute solution of carbonate stash. It forms obtuse double four-sided pyramids, which have a most se bitter taste. When acted on by ammonia in a close vessel, it dissipates, and after some time the solution becomes yellow, and deposits yelerystals, which are not bitter. When the ammoniacal solution is exto the air, a dark red substance is formed, which indicates an analogy sen this substance and orcine, which, as will be hereafter mentioned, in other lichens. Its composition is unknown, but it contains no nine. It is said to be powerfully febrifuge.

23. "Cetrarine is analogous to the preceding. It occurs in several 23, as in Iceland moss, Cetraria Islandica, and in Sticta pulmonacea. Extracted by alcohol. It forms a fine white powder, very bitter to the Concentrated hydrochloric acid colours it deep blue. Its other pro-

s are little known, but it is said to be used as a febrifuge in Italy.

24. "Elaterine is the active principle of elaterium, the inspissated of the fruit of Momordica elaterium. The elaterium is dissolved in Icohol, and the concentrated solution thrown into water, which precipithe elaterine. By repeating this process it is obtained pure. (Morries.) ms delicate silky crystals of a very bitter taste. One-sixteenth of a acts as a drastic purgative. Its composition is unknown. It merits re minute examination.

25. "Colocynthine.—The bitter and purgative principle of colocynth, is the pulp surrounding the seeds of Cucumis colocynthis. It is obtained by evaporating the infusion made with cold water, at first in oily which afterwards solidify into a brown, brittle mass. It is soluble tter, alcohol and ether, intensely bitter, and acts as a drastic purgates chemical characters are imperfectly known, and it is probably a

26. "Byronine.—Obtained by a somewhat similar process from the of the root of Byronia alba and B. dioica. It forms a brown or yela-white mass, having a taste at first sweetish, then acrid and very bit-boluble in water and alcohol, insoluble in ether. It appears to contain ren, and is probably a mixture of several compounds. It is a drastic tive, and has poisonous properties.

27. "Mudarine is found in the bark of the root of Calotropis Mu(Duncan.) It is soluble in water and alcohol. The aqueous solugelatinises when heated to 95°; at a higher temperature it is coaguthe mudarine separating as a viscid mass. On cooling, it is slowly
ampletely redissolved. Mudarine has powerful emetic properties.

28. "Scillitine.—Obtained from the juice of squills, the bulb of Scilla lima. A brittle mass, of a nauseous bitter taste. It acts as an emetic is a purgative, and appears to be poisonous. (Tilloy.)

29. "Cathartine.—Similar to the preceding. Obtained from the leaves tesia Senna and C. lanceolata, and from some other plants. It has a nauseous taste, and purgative properties.

• It forms greenish-yellow silky crystals, intensely bitter and astrinIt is very soluble in alcohol, and has neither an acid nor an alkaline
ion. Its action on the system has not been studied, but the bark is
as a remedy in the Antilles.

6531. "Columbiar.—Obtained from columba, the root of Menisperant palmatum. It is extracted by alcohol or ether. Forms colouries at transparent oblique rhombic prisms, or delicate white needles: a count fusible, and contains no nitrogen. It is very bitter, and becomes all and no when dissolved in acetic acid. It is the active principle of counts (Wittstock.)

When pure, it forms small white opaque prisms, which are intensely land and very soluble in alcohol. From the analysis of Wiggers, its forms is

probably Can Has Ot,

5538. "Lupuline is the bitter principle of hops, the female framed Humulus Iupulus. It is neutral, uncrystallizable, soluble in water solub-

cohol, and very bitter.

juice of Lactucase is the active principle of Lactucarium, the impossibilities of Lactuca sativa, L. virosa, and L. scariola. It forms yellow indistinct crystals, which have a strong persistent, bitter taste. It is pringly soluble in water, very soluble in alcohol. The anodyne effects lactucarium are most probably to be ascribed to lactucine.

5585. "Ergotine.—Discovered by Wiggers in the ergot of rye, & cornatum. It is obtained as a brown powder, of a pungent and bitter and and is conceived by Wiggers to be the active principle. He describe as narcotic and poisonous; but its composition and properties are unknown.

and it is most probably a mixture.

5536. "Porphyrozine.—Discovered by Merck in Rengal opina forms small brilliant crystals, which, when dissolved in diluted made toids and heated, yield a red colour. It is neutral, soluble in microbi ether, insoluble in water. It is quite distinct from the other crystales.

stances found in opium, but as yet has been but little examined.

sophila Struthium. It is extracted by alcohol, and purified by specialization from that solvent. It forms a white brittle mass, as the tallizable. It has a taste at first sweetish, then acrid and irritation, and the massive ancezing. It is soluble in water; and the solution, even when very the froths like a solution of soap. The root is used as a detergent.

hol from Sarsaparilla (Smilaz sarsaparilla). It is crystallumble, which in hot water and alcohol, colourless and tasteless. Its solutions have a property of frothing. Its formula appears to be C<sup>13</sup> H<sup>13</sup> O<sup>3</sup>. (Pograte Thubœuf; Petersen.) The Chinova bitter of Winkler, found in Chinacoa, has been shown by Buchner, jun., to be identical in its property of from that of smilacine only by 1 eq. of water.

Polygala senega and P. Sirginea. It is a white powder, at first takes afterwards very acrid, and causing a feeling of astringency in the gains. It also acts as a sternutatory. According to Quevenne, its formula

Cu Hu Ou,

6540. "Guaiacine.—Discovered by Trommsdorff in the wood and but of Guaiacum officinale. It forms a yellow britle mass, which has a simple acrid taste. It is no doubt one of the active principles of the guaranteed guaiacum, and is the cause of its acrid taste.

5541. "Plumbagine occurs in the root of Plumbago Europea. # #

tracted by ether, and forms fine orange-yellow crystals, which at first ve a sweet taste, followed by a burning acrid sensation. It is neutral, d soluble in hot water. Alkalies give to its solution a cherry-red colour, t acids restore the yellow. The root also contains a peculiar fat, not yet vestigated, which gives to the skin a lead-gray colour, whence the name the plant is derived.

5542. "Cyclamine: Syn. Arthanitine.—Found in the root of Cyclam Europæum. It crystallizes in fine white needles, of a burning acrid

ite, and having emetic and purgative properties.

moficinale. Extracted by alcohol. It forms delicate white prisms, fulle, insoluble in water, soluble in alcohol and ether. The solution has acrid burning taste. It is neutral. Formula, C<sup>4</sup> H<sup>5</sup> O. In some roots at had long been kept, Erdmann found a modification of peucedanine, difing from it only in being insoluble in ether. Its formula was C<sup>5</sup> H<sup>4</sup> O<sup>5</sup>; such only contains one atom of oxygen more than the formula of peucenine doubled, and was, therefore, probably formed from it by the action the atmosphere.

5544. "Imperatorine.—Found by Osann in the root of Imperatoria Osatium. Is extracted by ether. It forms long transparent prisms, has an rid burning taste, is neutral, fusible, insoluble in water, soluble in alcohol

d ether. Formula, C<sup>24</sup> H<sup>19</sup> O<sup>5</sup>. (F. Dæbereiner.)

5545. "Tanghinine.—Extracted by ether from the seeds of Tanghinia adagascariensis after the fixed oil has been removed by pressure. It is ystallizable; soluble in water, alcohol, and ether; very bitter and acrid.

is also poisonous. (Henry and Ollivier.)

5546. "Meconine.—Discovered by Couerbe in opium. It is dissolved, ong with most of the other ingredients of opium, when water is used as e solvent; and, being soluble in water, it remains dissolved when moria, narcotine, &c., are precipitated by ammonia. Part of it, however, is along with the precipitate. It is purified by the alternate action of alhol, water, and ether; in all of which it is soluble with the aid of heat. ben pure, it forms fine white prisms, which are at first tasteless, afterurds acrid. It is fusible, and may be sublimed unchanged. It requires r solution 266 parts of cold water, and 18 parts of boiling water. When ated with water, it first melts into an oily fluid, and gradually dissolves. ilphuric acid, diluted with half its weight of water, dissolves meconine, rming a colourless solution, which, when heated, becomes dark green. ater throws down from the green solution brown flocks, which dissolve alcohol with a rose-red colour. From this aicoholic solution the salts of rmina, lead, and tin, throw down fine red lakes. Meconine is quite neu-Its formula, according to Couerbe, is C<sup>10</sup> H<sup>5</sup> O<sup>4</sup>, or rather the half of s; but its composition cannot be considered as ascertained. By the aca of chlorine it is converted into mechloic acid, and nitric acid changes it to nitro-meconic acid.

5547. "Cubebine.—Found by Soubeiran and Capitaine in cubebs pepn (the seeds of Piper Cubeba). It is neutral, crystallizable, tasteless,
aringly soluble in water and alcohol. Its formula is probably C<sup>34</sup>
27 O<sup>30</sup>.

5548. "The following substances are neutral, have generally a bitter see or are tasteless, and are to a certain extent problematical, as the obveations regarding them are very imperfect. It is probable that many of em will be found identical with some of the preceding.

- "Alcornine, from Alcornico, the root of Hedwigia virgelioides.
- "Aliemine, from Aliema Plantago.
  "Arnicine, from Arnica montana.
- " Asclepine, from the root of Asclepias gigantea.
- "Absinthune, from the flowers of wormwood, Artemisia absisthing
- " Antiarine, from Antiaria toxicaria.
- "Amanitine, from Agaricus muscarius, A. bulbonis, and others.
- " Buenine, from the bark of Buena hexandra. " Canelline, from the bark of Canella alba.
- " Cascarilline, from the bark of Croton Eleutheria.
- " Cassiine, from Cassia fistula.
- " Centaurine, from Erythraa Centaurium.
- " Colletine, from Colletia spinosa.
- " Coriarine, from Coriaria myrtifolia.
- " Cornine, from the bark of the root of Cornus florida.
- "Corticine, from the bark of Populus tremula. "Cytistne, from the seeds of Cytisus Laburnum.
- "Daphnine, from the bark of Daphne Mezereum and other species is crystallizable.
  - "Datiscine, from Datisca cannabina.
  - " Diosmine, from the leaves of Diosma crenata.
  - " Euonymine, from the seeds of Euonymus Europana.
  - "Fagine, from Fagus sylvatica.
  - " Frazinine, from the bark of Frazinus excelsior.
  - "Geraniine, from the Geraniacea.
  - "Granatine, from unripe Pomegranates.
  - "Guacine, from Guaco leaves.
  - "Hesperidine, from the spongy part of the Orange rind. Crystalli
  - " Hyssopine, from Hyssopus officinalis.
  - " Ricine, from Hex aquifolium. Crystallizable.
  - "Lapathine, from Rumez obtusifolius.
  - "Ligustrine, from the bark of Ligustrum oulgare.
  - "Lilacine, from Syringa or Lilac.
  - "Liriodendrine, from the bark of the root of Liriodendron twiisifa"
  - "Menyanthine, from Menyanthes trifoliata.
  - "Melampyrine, from Melampyrum nemorosum.
  - "Narcitine, from Narcissus pseudo-narcissus.
  - "Olivile, from Olea Europæa.
  - "Olivine, from the leaves of Olea Europæa.
  - "Primuline, from the root of Primula veris.
  - "Pyrethrine, from the root of Anthemis Pyrethrum.
  - " Populine, from the bark and leaves of Populus tremula.
  - " Phillyrine, from the bark of Phillyrea media and latifolia.
  - "Rhamnine, from Rhamnus frangula.
  - "Scordine, from Teucrium Scordium.
  - " Scutellarine, from Scutellaria lateriflora.
  - "Serpentarine, from Aristolochia serpentaria.
  - "Spartime, from Spartium monospermum.
  - "Spigeline, from the root and leaves of Spigelia anthelmia.
  - " Tanacetine, from Tanacetum vulgare.
  - " Tremelline, from Tremella mesentherica.
  - "Zedoarine, from the root of Curcuma aromatica."

#### ETHERS, AND THEIR COMPOUNDS AND DERIVATIVES.

# Of Ethyl Ethers (3090).

549. Agreeably to the classification proposed in treatof ethyl (3077), common ether, the oxide of that comid radical, as the first in the class of simple ethyl rs, is primarily to be the object of attention.

# the Oxide of Ethyl, common Ether, erroneously called Sulphuric Ether, C<sup>4</sup> H<sup>5</sup>O.

550. It has been mentioned, that this compound is now ed ether, on account of a sort of prescriptive claim, aligh the name by which it is designated has been apriated to a class of bodies having, in common with it, e important characteristics (3083).

551. Of the Properties of the Oxide of Ethyl.—The oxof ethyl is a colourless, transparent, volatile liquid, ng only seven-tenths of the density of water, or seven-

ths of that of absolute alcohol.

552. It is so inflammable, that a jet of it may be ined throughout its whole length, when extending many

It has a fragrant smell and an aromatic taste, which, ough pungent and stimulating, is not unpleasant. ity to that of water at 60°, is as 725 to 1000. It boils reen 97° and 98°, and congeals before it reaches the perature of — 47°. With alcohol it unites in all proions, but may be recovered therefrom by agitation twice its bulk of water, which, combining with the hol, subsides gradually, allowing the liberated ether to

1 a superstratum easily separable.

553. One part of ether dissolves in ten of water, and part of this liquid in thirty-six of ether. Essential oils soluble in ether to any extent, and also the margarine olein of fixed oils; but stearine is so little soluble in er, that it is employed to depurate it of the two other we mentioned constituents of fat. Ether is likewise a vent of most of the resins. It takes gold, in the metalstate, from a solution of the chloride of that metal, ning an ethereal solution, which has been employed to steel. It dissolves several of the haloid compounds, cially the chloride of zinc and bichloride of mercury;

also several organic acids, the acetic, gallic, benzoic, oleic, and stearic, for instance. The solubility of various as ganic alkalies in ether has been mentioned in treating of their extraction. Of sulphur, it takes up 1/2 of its weight; of phosphorus, from 1/2 to 1/2, according as it is more or less free from water. Bromine and iodine are coposity soluble in ether, the solutions being, however, habit to spontaneous decomposition, producing bromohydric and iodohydric acid, and some other products which have not been studied.

ately, each bubble inflaming spontaneously at the ordinary temperature of the air, giving birth to chlorohydric acid, and liberating curbonic scid. As hydrous sulphuric acid, in the cold, generates from other, isotheric, and ethionic acid, besides heavy oil of wine, light oil of wine (5537), and so phovinic acid. At a high temperature, these acids are resolved into heavy oil of wine, water, other, sulphurous acid, and olefant gas.

5555. Nitric acid, aided by heat, converts ether into formic, oxalic, and

carbonic acid, together with aldchyde.

5556. Of chlorohydric acid gas, ether absorbs a large quantity;

by distilling a concentrated solution, chloride of ethyl is generated.

5557. Dry alkaline hydrates have no reaction with pure other a start nary temperatures, but when moisture and oxygen are present and best employed, cause it to become brown after some time, and to form after acetates or formiates. Potassium and sodium are alleged, by Loobs, story to deoxidize other, and finally to decompose it into gaseous and only care hydrogens, forming oxides with which the other combines as an accident

5559. In presence of iron, lead, or zinc, with access of oxygen, the rie

ment is absorbed, generating acetates.

5559. Ammoniated ether may be obtained by subjecting ether, shied

lime, and chloride of ammonium, to the distillatory process.

5560. An etherial solution of the bi-todide of mercury is obtained by the solution of one part of the bi-todide in twelve parts of other. On agreement of the bi-thloride of iron dissolves in four parts of other. On agreement aqueous solution of this bichloride with other, this liquid takes the bithloride from the water, forming a golden-yellow liquor, from which light causes crystalline protochloride to precipitate.

5561. In consequence of the solubility of narcotina, and insolubility of morphia in other, it is employed to denarcotize opium in preparing s for

making denarcotized laudenum.

5562. The tension of the vapour of other being, per sc, adequate to port a pressure about half as great as that of the atmosphere, it computes the volume of any gas to which it may be added. The may be made evident by introducing a measured quantity of any gas is volumescope, and adding, subsequently, a portion of ether. (818).

Agreeably to the experiments of Dalton, the vapour of any liquid in contact the air, or any permanent gas, supports a proportion of the atmospheric pressure, which bears the same ratio to the whole pressure, as the height of the column of the cury, which the vapour in question will support, per ee, in an exhausted receive.

of that gas was mingled with four volumes of hydrogen and two of oxygen, no condensation ensued when the mixture was ignited. The elements of the gas combining with those of water in the act of uniting, generated a new gas containing the elements of both. It was likewise mentioned, that half a volume of ether had about as much efficacy when substituted for the olefant gas, as a whole volume of the latter.

5564. This is interesting, as tending to show that in the same space ether vapour contains about twice as much carbon and hydrogen as olefiant gas

(1276).

5565. Of the Means of obtaining Ether.—Respecting the means by which ether is elaborated, a general explanation has already been given in treating of ethyl (3804). The old recipe for its manufacture, was to distil two measures of officinal alcohol of about 0.840 with one of sulphuric acid, without any subsequent addition of alcohol; but, latterly, the proportions have been nearly reversed by using, at the outset, nine parts of acid, by weight, with five parts of alcohol, the proportion of this liquid being sustained by subsequent additions, compensating the diminution resulting from the vaporization of the products. Liebig recommends, that in using these proportions, the alcohol be added to the acid in a copper or cast iron vessel, the liquids being mingled by stirring them with an iron spatula; but agreeably to the experience of a manufacturer in this vicinity, who, for many years, was in the practice of distilling a large quantity of the etherifying materials at a time, it is preserable to introduce the alcohol into the alembic first, and then the acid, in a continued stream. This stream, by its superior weight, produces a descending current, carrying along with it the alcobol with which it comes into contact, and forming a compound, of which the boiling point is about 280°. The descending current displacing the liquid previously near the bottom of the alembic, causes it to ascend at the sides, and thus establishes a circulation, by which a complete intermixture The heat generated meanwhile, acting upon If the materials is effected. ome of the alcohol not in contact with the acid, is, in a greater or less deree, expended in vaporizing a portion of this ingredient, which, condensing the receiver, should be restored to the body of the still or retort employed. 'his method of manipulation, to which I have myself long resorted, has seeral advantages over that of Liebig; agreeably to which, the alcohol being pured over the acid, and in contact with the air, must sustain some loss by The mixture being made in one vessel, and the distillation in nother, causes unnecessary trouble, and the heat generated by combinaon is lost, which, in the other case, requires little aid from the fire emoyed to cause the distillation to commence.

5566. The most advantageous method of applying heat, in this and many her cases, is that already described of a furnace having coals in a drawer hich can be withdrawn in an instant, partially or wholly, so as to render temperature perfectly controllable (963). Where carburetted hydrogen

the height of the mercury in the barometer at the same time. Hence, as the common which the vapour of ether will support, per se, at ordinary temperatures, is post half that of the usual height of the barometric column, it follows, that when said ether is introduced into any gas, its vapour relieves the gas of half the pressure, at the same time deprives it of half the space, so that they require twice as much som as the gas required, per se; consequently the volume of the mixture becomes since as great as that of the gas previously.

is supplied to a laboratory, from a gas light establishment, and a glass :tort is to be used, a tube, forming a circle of four or five inches diameter. and perforated at intervals of about half an inch, so as to allow in its currenference from twelve to twenty gas lights, forms an efficient mean of appying a competent and manageable heat. If the distribution of the ethenium materials be carried on until the resulting carbonaceous mass swells up as as to endanger its coming over, it will be found that the first product comsist of ether with undecomposed alcohol, then other and water, and afterwards ether with sulphurous acid and heavy oil of wine, forming a wine liquid. But, according to Liebig, if before alcohol ceases to come come minute proportion, absolute alcohol be gradually added by a tube, was a very small aperture at the lower end, terminating under the surface of the mixture so as to keep it at the same level, by compensating the damage. resulting from the distillation, the evolution of ether and water contents without the extrication of sulphurous acid and oil of wine. If, says Lebes "the operation be well managed, only ether and water will be evolved, and the acid may serve for the preparation of ether, indefinitely, without per ceptible diminution." When alcohol of the officinal density is used, in the way thus proposed, the acid soon becomes too much diluted to person the office of an etherifyer. Liebig admits that when the spurit of wine copier ed, contains 90 per cent. of anhydrous alcohol, only 31 parts can etherified by 90 of sulphuric acid, and when the proportion of water to in acid exceeds the ratio of 9 to 2, ether cannot be evolved."

have been accustomed to employ an inverted open-necked bell-glass, through the axis of which a glass tube passes, being made to form an anti-tight passes as to embrace the neck, by means of a gum elastic bag, cut off near the brain so as to embrace the neck of the bell, while its own neck embraces the take being secured to both by ligatures. The beak of the retort is drawn by means of a fire, and bent at right angles so as to descend into the passes of the refrigerating tube. The bell is supplied with icr and water, this liquid being drawn off by a syphon as the ice melts, in order to a more to be added. The refrigerating tube must terminate with a bottle, surrounded by ice and water. It is usually recommended to appear the othereal portion of the product, and rectify it over milk of that, at a

That sulphovinic acid is the inevitable consequence of mixing and hester phuric acid with alcohol beyond a certain point, has already been mentioned is the ing of ethyl (3036) and of sulphovinic acid (5297). This combination arrows proving from the affinity of hydrous sulphuric acid, sulphate of water, when undistributed etherine, and for more water, so that while one portion attracts the other of the about the other attracts the water.

Agreeably to the representations of Liebig, above stated, if, during the ethicition of alcohol by sulphuric acid, that ingredient be supplied in the proportion vary to compensate the evolution of other and water, a given quantity of acid varies stances, only the alcohol appeared to undergo decomposition, being noticed to cherlith, led him to infer that the part performed by the acid was merely extended for infer that the part performed by the acid was merely extended for aulphovinic acid whenever alcohol and sulphuric acid are mingled in deeperformed for aulphovinic acid whenever alcohol and sulphuric acid are mingled in deeperformed for aulphovinic acid present, and yet absorb alcohol to form another portion of the sulphovinic acid present, and yet absorb alcohol to form another portion of the said. The most feasible explanation is, that the contact of the alcohol with the said. The most feasible explanation is, that the contact of the alcohol with the said can combine with exide of ethyl, while in other parts of the mixture temperature may be sufficiently high to cause other atoms of the mane has dismaited from the acid.

caustic alkaline solution, with the heat of a water bath of about 120°. I have found ammonia the most speedy agent for this depurating process.

5568. Agreeably to the old process, oil of wine was generated towards the last. Hence, after the ether was distilled, a compound of alcohol and oil of wine remained, and could be brought over by raising the water bath to a boiling heat. Hoffman's anodyne liquor was thus obtained.

Of heavy Oil of Wine, denominated by Liebig, "the double Sulphate of the Oxide of Ethyl and Etherole," the true Sulphuric Ether, C<sup>4</sup> H<sup>5</sup> Q + C<sup>4</sup> H<sup>4</sup> + 2SO<sup>3</sup>: also of light Oil of Wine.

and alcohol employed to produce ether, becomes sufficient to retain the ether until the temperature rises above 324°, a reaction ensues by which a yellow, sulphurous, ethereal solution of oil of wine comes over (3039) (5299). This consists of nearly equal parts of sulphurous acid and ether, the oil of wine being present only in a comparatively minute proportion. This liquid being subjected to distillation at a heat not exceeding 120°, the greater part of the ether and sulphurous acid may be brought over. The residue may then be exposed in vacuo over sulphuric acid and slaked lime. By these means all the sulphurous acid ether, and water, are absorbed, the oil of wine being isolated.

5570. Properties.—Thus obtained, oil of wine has an unctuous consistency, whence its name. It is transparent, nearly colourless, and highly

fragrant. Its taste has a resemblance to that of peppermint.

Hennel, and afterwards more fully by Serallas, that kind of oil of wine which is designated as "heavy," was shown to be a chemical compound of sulphuric acid, carbon, and hydrogen. Subsequently, it was considered as a neutral hydrated sulphate of etherine, 2C<sup>4</sup> H<sup>4</sup> + HO + 2SO<sup>2</sup>. This, of course, contains the same elements as if it were considered as an anhydrous seutral sulphate of the oxide of ethyl, 2C<sup>4</sup> H<sup>2</sup>O + 2SO<sup>2</sup>. Lately, it has seen represented by Liebig, as a double sulphate of the oxide of ethyl and therole; this last mentioned ingredient being, in other words, etherine, C<sup>4</sup> H<sup>4</sup>.

5572. Oil of wine, thus defined, has been called heavy oil of wine, because it sinks in water. It appears that it may be more or less deprived of the sulphuric acid, by being distilled from milk of lime, or by being digested with caustic alkaline solutions, and then forms what is called light oil of wine, being lighter than water. From the heavy oil, when free from water, I was unable to remove the acid entirely by distillation from potassium.

5573. When alcohol is etherified by chloride of zinc, two light oils are alleged to be evolved, one having the formula C<sup>8</sup> H<sup>7</sup>, the other C<sup>6</sup> H<sup>9</sup>. Being devoid of sulphuric acid, these oils are of course quite different from the keavy oil, of which the formula is above given. The allegations respecting the composition of this heavy oil, are to me quite unsatisfactory, and lead to the impression that we are still ignorant of its true constitution. Nothing can be more anomalous, and inconsistent with the laws of chemical combination, with which experience has made us acquainted, than that two atoms of an acid, being comprised within a compound, and one of them in union with an oxidized radical acting as a base, as ether does, the other should refuse to unite with another atom of that base, and yet combine with a non-pridized radical, etherole or etherine. In its free state, this last mentioned compound unites neither with sulphuric acid, nor any other acid, and yet

it is represented as replacing the basic water, and completely neutralized the acid properties of sulphovinic acid, so that no immediate reaction such as on contact with the most powerful bases. It is unnecessary to such here the suggestions respecting the bibasic character of sulphovinic ad-

made in treating of its mexplicable properties (5289).

5574. Of Hoffman's Anodyne Liquor.—In consequence of the many tions made in the manufacture of ether, with the view of saving the soci agreeably to the explanations above given (5566), the genuine another is quor of Hoffman, being no longer a collateral product of that management a mixture of ether and alcohol came to be substituted in commerce is the true medicine. This drew the attention of some of our older physics. Dr. Wistar, and my late colleague, Dr. Physick. Dr. Wistar had meaded that the modern anodyne liquor did not produce any mulkiness in water when added to it, and he observed that the presence of this appeared o sential to the efficacy of the medicament. In consequence of the report of Dr. Physick, having given some attention to the subject, I ascertaint that in the officinal anodyne there was generally no oil of wine, and bas nothing to be separated on the addition of water. This phenomeron was found only to ensue in the anodyne prepared by those druggists who is bered to the old method of manufacture. As both by Drs. Physick of Dewees, much value was attached to the real anodyne " as highly wife in some disturbed states of the system, in tranquillizing and disposar sleep," I regret that no efforts have been made, by those who are a the practice of medicine, to ascertain whether there is any separate efficient the oil of wine, or whether it operates by giving greater permanency is the impression made by ether by lessening its volatility; and if this he the man whether other essential oils cannot be used in heu of oil of wine, as a \* hicle for ether.

tioned, that when the materials employed for the generation of eiter has a certain ratio, and the temperature reaches a certain height, a velice to comes over, which consists of heavy oil of wine, ether and sulphuranted. This liquid being refrigerated by ice, and mingled, gradually, was anothing, also refrigerated in a bottle surrounded by ice water, the ethered tion loses about half its bulk and weight. The residual liquid, which have upon the resulting ammoniacal solution, being separated by dilution with twenty-four parts of alcohol, forms the anodyne liquor which I have been toned.

accustomed to prepare.

#### Of Alcohol, or the Hydrated Oxide of Ethyl.

of alcohol was, I trust, sufficiently explained (3069). I have now to treat of the means of obtaining it, and of the

properties.

5577. Alcohol can only be obtained through the medium of the process called the vinous fermentation, being that by which the juice of the grape, of the apple, or pressor infusions of sugar, or farinaceous substances, are repetered spirituous. By subjecting fermented liquors the originating, to distillation, alcohol, diluted with water, and

flavoured by various peculiar empyreumatic oils, is obtained, being known as brandy, rum, or whiskey, accordingly as it may be derived from wine, from molasses, or

from grain or cider.

5578. The vinous fermentation may be induced by the addition of yeast to a solution of sugar, kept between 60° and 70°. During this process, a new distribution of the elements takes place, so as to form alcohol and carbonic acid. One atom of dry grape sugar, C<sup>12</sup> O<sup>12</sup> H<sup>12</sup>, is converted into two atoms of alcohol, 2(C<sup>4</sup> H<sup>5</sup> O + HO) and four atoms of carbonic acid, 4CO<sup>2</sup>.

Two atoms of alcohol, - - C<sup>8</sup> H<sup>19</sup> O<sup>4</sup>
With four atoms of carbonic acid, C<sup>4</sup> O<sup>8</sup>

Form one atom of sugar, - C<sup>19</sup> H<sup>19</sup> O<sup>19</sup>

5579. It can hardly be necessary to mention, that the intoxicating power of the various liquids known generally in commerce as spirits, as well as that of wine, beer, cider, and other fermented liquors, is due to the alcohol which they contain. These spirits, whether known as whiskey, gin, rum, brandy, or arrack, in a chemical point of view, may be considered as mixtures of water with alcohol. Proof spirits is a term applied to any of these mixtures, when consisting of their principal ingredients in

equal proportion.

5580. When, in consequence of the request of the British treasury department, a committee of the Royal Society undertook to make a table, showing the relation between the density and the quantity of alcohol in a series of mixtures of this liquid and water, though the most scrupulous accuracy was displayed, the conclusion was adopted, that the matter existing in the various kinds of spirit, on which their diversity as respects flavour and value is dependent, was too small to require to be taken into account. Nevertheless, it is well known that peculiar volatile oils accompany the whiskey obtained from grain and potatoes; and Ure alleges, that spirit obtained from damaged grain, has been found to contain a peculiar volatile matter augmenting its intoxicating power, so as to produce in some persons a sort of frenzy. This matter, at the end of a few months, was spontaneously decomposed, so as to render the spirit less nauseous and unwholesome. The impression which has existed in this country, that brandy is more unwholesome than other spirituous le may depend on an analogous cause. I am under the pression that brandy and rum contain principles cause their peculiar flavour, and that the difference tween old and new spirit, is due to the modification those essential oils on which the peculiarity of quantum principles.

in such cases dependent.

5581. By distilling one-half of the volume from area spirit, officinal spirit of wine is procured, and by o rectification, a liquid of the density 0.825 may be obtain still containing eleven per cent. of water. But it is it as it sible for the vapour of any liquid to be formed in the !! sence of another liquid, without becoming associates a portion of its vapour. Besides, the inferior dense aqueous vapour creates in it a tendency to rise with vapour of alcohol, as hydrogen does in atmospher Hence the presence of two or three per cent. of was is alleged, makes the boiling point of alcohol lower. sequently, a more aqueous portion distils first under circumstances. But on the other hand, when the pretion of water reaches to six per cent., the result is us ed, so that the product, which first comes over, is aqueous than the subsequent product. According to ning, if the capital of the still be kept at 174°, no which contains less than ninety per cent. of alcohol too pass over. Of course, the same object would be obtained by passing the beak of one retort into the tubulure of 300 ther quite empty, and preserving the latter at a protemperature, while its beak is made to communicate 🖘 a receiver properly refrigerated.

5582. Alcohol may likewise be concentrated by subjected, in a well cleansed bladder, to the temperator of 122°. The bladder is made more efficient by smeared with a solution of gelatine, four times inside

twice outside.

5583. But to procure absolute alcohol, or, in words, that which is devoid of water, a resort must be to a chemical agent having a great affinity for water. cently ignited carbonate of potash, quick-lime, or chloride of calcium, may be employed. In either case, spirit must be kept in contact with the substance employed for some time before distillation. Chloride of calcium.

y fused, is generally preferred. Of the spirit of wine at more than 0.833 in density, and of the chloride of ann, equal weights being mixed so as to form a satusolution by the distillation of this and a well consolution to the configuration of the configuration of the spirit of wine as the configuration of the spirit of wine as the configuration of the spirit of wine as the configuration of the spirit of wine at the configuration of the spirit of wine as the configuration of the spirit of wine as the configuration of the chloride of the chloride of the chloride of the spirit of the chloride of the c

ding to Ure, of 0.791 at 68°. 34. Alcohol has a very powerful affinity for water, so absorb it from the atmosphere, and from organic ances in general. It is by neutralizing water that it rves anatomical preparations, performing, in this re-, a part analogous to that of brine. As the freezing of mixtures of this liquid with water is extremely low added to snow, it operates as deliquescent salt, and aces cold (419). The opposite effect results from its with water, as it forms in that case a liquid, of which mpacity for heat is less than the sum of the capacities ingredients. Alcohol, by combustion, yields only and carbonic acid. It is more expansible than wand boils at 176°. Its capacity for heat, whether in equid or aëriform state, is much less than that of It is a powerful solvent, and highly useful agent armacy, and in the delicate analysis of vegetable and al matter. There is no satisfactory evidence that al-has ever been frozen. The most intense cold proby solid carbonic acid and ether, by Dr. Mitchell, ed it to become syrupy in consistence, but did not it. The addition of one-seventh of oil of turpentine ender the flame of alcohol so luminous, as to be a etent substitute for a candle flame. When alcohol sed through a red-hot porcelain or copper tube, it is mposed into water and carburetted hydrogen.

# Of Ethero-sulphurous Acid, or Sulphurous Ether.

5. Although no definite compound of sulphurous acid with the oxide pl has been made, an affinity exists between this acid and oxide, reag that between alcohol and water. Sulphurous acid boils at —12°, at 98°, the difference being 110°. Of course, were not the affinity in these fluids more energetic than that between alcohol and water, of the boiling point differs only by 36°, they would not remain united at ry temperatures. The boiling point of sulphurous ether is lowered, portion as the ratio of the acid to the sulphuric ether is increased, it contains oil of wine, the temperature necessary to ebullition of the rate, is inversely as the quantity of the sulphurous acid, and directly tof the oil of wine, to the quantity of the other ingredient. Hence,

although I have obtained sulphurous ether, which boils at 26°, it is not possible, with the heat of a boiling water bath, to separate the last portion of this ether from the oil of wine, since a part of the latter distributed with at I kept twenty-six measures of the compound of sulphurous ether and of wine in a glass measure, over water, for three weeks, without the segurity perceptible diminution of the quantity of the former. By means of a separate percent by acrews, about an ounce of the volatile sulphurous ether will kept in contact with water for more than six weeks without apparent with the ether to the alkali takes place slowly, unless agitation be employed.

5586. Of Hyponitrite of the Oxide of Ethyl, Hyponitrie of Ethyl, Nitrite of Ethyl, Nitric Ether, Nitrons Ethes, C' HO' + NO'. This ethereal compound is generated by the mixture of alcohol with nitric acid, provided the comcentration and proportion of the latter and the temperature, be such as to prevent the reaction from being to violent; in which case the products are liable, according to Liebig, to be carbonic, acetic, and formic acid, with acetic and formic ether. This distinguished chemist ond to mention the residual elements of the nitric acid employed. From the copious display of red fumes, there seems to be reason to infer that nitrous or hyponitres acid is abundantly evolved. It is alleged by the same author, that when the reaction is sufficiently mitigated by the dilution of the reagents, and moderation of the teaperature, only aldehyde and hyponitrous ether are generated.

5587. Of this I presume the following rationale may be given:—From an atom of the acid employed, two atoms of oxygen, uniting with two of the hydrogen of an atom of the alcohol, convert it into aldehyde. Meanwhile three atoms of oxygen, remaining united with one of nitrogen. the state of hyponitrous acid, combine with an atom of the oxide of ethyl, expelling the water by which it was enabled to exist as alcohol. It follows, that at a minimum, one half of the alcohol must be destroyed.

5588. According to Liebig, the best process for the generation of this ether, in purity, is to impregnate alcohol with the vapour resulting from the reaction of nitric activity starch, passing the nëriform proceeds through a well-refrigerated tube to a receiver in a similar state. I have repeated this process twice, and have found a very small quantity of pure ether to be produced, with comparatively large consumption of the materials.

5589. I conceive that the best process is that of which I gave an account about four years ago, and which is as follows:—

5590. Fourteen parts of the hyponitrite of soda with just enough water for its solution, seven parts of alcohol, eight of sulphuric acid diluted with twelve parts of water, are to be refrigerated, and introduced into a bottle immersed completely in water. In a very short time, hyponitrous ether will be seen swimming on the mixture; and after about six hours the process will be so far perfected, as to make it expedient to decant the ether.

5591. In lieu of including the materials within a bottle, as above described, the salt, previously dissolved in water, may be introduced into a pubulated retort, with a beak recurved and adapted to a refrigerating apparatus and receiver surrounded by ice-water, as already described. Through the tubulure of the retort, a tapering glass tube, terminating in an orifice of about a tenth of an inch in diameter, should descend nearly to the bottom, being secured air-tight to the tubulure by gum elastic or other lutings.

5592. The alcohol, acid, and water, being united and quite cool, may now be poured in through the tube; the ether rapidly generated is condensed in the receiver in a state quite free from aldehyde. Water containing a very little lime, potash, soda, or ammonia, may be used to free it en-

firely from acid, and quick-lime to free it from water.

5593. Hyponitrous ether, thus obtained, differs from the ether ordinarily known as nitric or nitrous ether, in having a more bland and saccharine taste, milder odour, and greater volatility. It boils below 65° F., and, by its spontaneous evaporation from the bulb of a thermometer, produces a cold of 15° below zero, F. Touched with the finger or tongue, it hisses as does water with a red-hot iron.

5594. If, after having boiled for some time, it be allowed to stand for while at a temperature below its boiling point, the boiling will recommence at a lower temperature than that which was indicated by the thermometer

when the boiling ceased.

an ethereal gas, which appears to be formed by the materials by which the liquid ether is generated, even when refrigerated below the freezing point. I have collected this aëriform ether, in large quantities, in bells over mercury. When subjected to great pressure, it condenses, more or less, into a yellow liquid, which produces, when allowed to escape into the mouth or mostrils, the same impression as the liquid ether. I have conjectured that this ether might be a compound of the liquid ether with nitric oxide gas, or that it may be isomeric with the liquid ether. Notwithstanding many efforts to obtain a liquid ether not resolvable partially into this gas, I have never succeeded. Hence the boiling point is extremely variable, as I have seen bubbles escaping below 40° from the liquid ether, when recently condensed after distillation.

5596. In the production of cold by mixture with solid carbonic acid, Dr. J. K. Mitchell found this ether more efficacious than that commonly known

sulphuric ether, more properly called hydric ether.

5597. When the new ether, as it is first evolved, is distilled from powdered quick-lime, this earth imbibes an essential oil, which, with the aid of senter, is yielded to pure hydric ether. Of course, it is easy to remove this solvent by evaporation or distillation. 5598. The odour of this oil seems to be an ingredient in that of nitric other. Possibly the hyponitrous ether may resolve uself into this oil and the gaseous ether, so that its boiling point may be varied by this chemical change. I suspect that the essential oil in is one of the impurities which causes the boiling point of the other; by nitric acid and alcohol, to be higher than the boiling point of

tained, as in my process, by nascent hyponitrous acid-

5599. When the heat is raised, after the volatile other crases over from the materials above mentioned as producing it, othereal are distilled, of which the boiling point gradually rises as the proceeds. Meanwhile, the product thus obtained, becomes more acrid, till at last it is rendered insupportable to the tongue, as relater tasts. On mingling these liquids with a solution of green a tron, the ether is all absorbed; but an acrid liquid, which cause tasts, is not absorbed, and may be separated by hydric other. being vaporized by heat, the acrid liquid remains. The smalle this liquid is productive of an effect upon the organs of tasts and that of mustard or horse-radish.

5600, The new ether, when secured in a glass phial by means ground stopper, does not undergo any change by keeping in a cot for several months. A phial was suspended about fifteen feat surface of the ground, in a cistern of water, for about five months was left in a cool cellar for a longer period, without any appara of properties. In this case pressure prevented the escape of the ground.

as above mentioned.

5801. All the ethereal compounds formed by the reaction of to of nitrogen with alcohol appear to be decomposable by green iton. Under these circumstances, according to Berzelius, a male

is formed from common nitric ether-

alcohol, and liberates nitric oxide gas, which is, it is well knot absorbable by the green sulphate above mentioned. Let there be lindrical glass jars, of such a ratio to each other in size, as to all terstices of about half an inch between the second or intermediate the outer and innermost jar; likewise, let two bell-glasses be pushed a size as that one of them may enter the inner interstice, other will cover and descend into the outer interstice. Let a u containing the other, be placed in the innermost jar, and let the o supplied with green sulphate of iron, the other two with concentration, and let the bells be put in their respective places.

5603. Under these circumstances, the ether will be gradually and the alcoholic elements, with some oxygen, will be absorbed b while nitric oxide, being liberated, will pass into the sulphate, and

sequently, absorbed.

#### Of the Process for Sweet Spirit of Nitre.

5604. This name is applied to a dilute solution pure hyponitrous ether in alcohol, which has acquamme from being obtained by subjecting nitre and ric acid to distillation with a great excess of alcoholoproportions, agreeably to the United States Disper

ounds of nitre, one and a half pounds of acid, nine of alcohol, the product being rectified from a of spirit and an ounce of carbonate of potassa. t spirit of nitre of commerce is a very uncertain to the nature and proportion of its ingredients, been informed by eminent druggists, as well as By keeping, it becomes partially acidified, have kept pure hyponitrous ether in a cool celearly a year without deterioration. I am of opiit would be advantageous if the prescriptions of cians were made with reference to ingredients of gree of purity. The physician should know how I ether is contained in the diluted article which his patient to use. Hence the pure hyponitrite of ethyl should be prescribed, adding as much alvater as may be deemed necessary. Agreeably sent practice, it is in the power of manufacturing to impoverish ethereal preparations with little detection.

Pursuant to the London Pharmacopæia, three nitric acid, by distillation with a quart of alcoillowed to produce twenty-four fluid ounces of it of nitre. According to Thenard, the quantity when the materials are in the ratio of equality, o two-thirds the weight of the acid. Hence it is that the quantity of ether in twenty-four fluid sweet spirit of nitre, obtained as above mennot more than two ounces. I infer that sweet nitre, of a more uniform strength, would be obthe addition of alcohol to pure nitric ether, to an more than adequate to render it soluble in water, adding water to the alcoholic solution, until the uld form only a twelfth of the aggregate. In a on thus made, the properties of the ether would necessarily associated with those of alcohol, as al officinal preparation.

Perchlorate of the Oxide of Ethyl, or Perchloric Ether.

is ether was discovered, in my laboratory, by Mr. Martin Boyérk Hare.

was obtained by subjecting about ninety grains of crystallized of baryta, with an equivalent proportion of perchlorate of badistillatory process, receiving the product in from one to two absolute alcohol. By complex affinity, the sulphuric acid of the

sulphovinate dispossesses the perchloric acid of the baryta, while, at the same time, the last mentioned acid combines with the oxide of ethyl.

5608. The perchlorate of ethyl is a transparent, colourless house, posessing a peculiar, though agreeable smell, and a very sweet taste, which on subsiding, leaves a biting impression on the tongue, resembling that of the oil of cinnamon, but more acrid and enduring. It is between that water, through which it rapidly sinks. It explodes by ignition, friction, or serve sion, and sometimes without any assignable cause. Its explosive pagertal may be safely shown, by pouring a small portion of the alcoholic with into a small porcelain capsule, and adding an equal volume of water. The ether will collect in a drop at the bottom, and may be subsequently as rated by pouring off the greater part of the water, and throwing the rest a moistened filter, supported by a wire. After the water has dra med 4 the drop of ether remaining at the bottom of the filter may be explain either by approaching it to an ignited body, or by the blow of a bases. The violence and readmess with which this ether explodes is not surposed by that of any other known compound. By the smallest drop, so and porcelain plate may be reduced into fragments, and by a larger quantity. powder. In consequence of the force with which it projects the man fragments of any containing vessel in which it explodes, it is necessary we the operator should wear gloves, and a close mask, furnished with the glass-plates at the apertures for the eyes, and perform his manufacture with the intervention of a moveable wooden screen.\*

5609. In common with other ethers, the perchlorate of ethyl is inside in water, but soluble in alcohol; and its solution in the latter, when we ciently dilute, burns entirely away without explosion. It may be kept to a length of time unchanged, even when in contact with water; but the station of this fluid, when employed to precipitate it from its alcoholic causes it partially to be decomposed. Potassa, dissolved in alcohol, at added to the alcoholic solution, produces immediately, an abundant produce of the perchlorate of that base, and, when added in sufficient quarter.

decomposes the ether entirely.

5610. The perchlorate of ethyl has been subjected to the heat of balls.

water without explosion or ebullition.

bination of an inorganic acid containing more than three atoms of cases with the oxide of ethule, and that the chiorine and oxygen in the wind compound are just sufficient to form chlorohydric acid, water and carbon oxide with the hydrogen and carbon. It is also the only ether which are plosive per se.

#### Of Acetic Ether, or Acetated Oxide of Ethyl, Ca Ho Oa + Ca HO.

5612. In common with other oxacid ethers, this ether may be obtained agreeably to the principles already set forth (5303), by distilling alcoholand sulphuric acid, or in other words, sulphovinic acid, with any access, of any sulphovinate with concentrated acetic acid.

5613. Acetic ether is colourless, burns readily with a pale vellow has a refreshing odour, with a density of 0.890 at 60°. It boils at 16°, does not redden littuus, is soluble in seven parts of water, and to every pro-

<sup>\*</sup> For the particulars of the process I refer to the American Philosophical Transtions, Vol. 8, New Series; also to Silliman's Joseph, Vol. 42, for 18th, page 62

on in alcohol or ether. In general it is a solvent of all the substances h dissolve in this last mentioned liquid. By alkalies it is readily mposed, likewise by sulphuric acid by which it is resolved into ether acetic acid.

#### f Oxalic Ether, or Oxalated Oxide of Ethyl, C4 H5O + O4 H3 O3.

314. This ethereal compound, discovered by Thenard, may be obtained he following process:—Four parts of binoxalate of potash are mixed in ort, with five parts of oil of vitriol, and four parts of alcohol, of 840°, briskly distilled. As soon as the product becomes turbid on the addiof water, the receiver is changed. The subsequent product being kly mixed with four times its bulk of water, the ether sinks to the bot-It should be separated and washed with successive portions of water, t becomes neutral to test paper. The ether thus washed is transferred small dry retort, filled up to nine-tenths of its capacity, and rectified. oon as the product becomes clear, and the boiling goes on, regularly, receiver is changed. What now passes over is pure anlightous oxalate re oxide of ethyl (oxalic ether)—(Ettling). It is a colourless, transpaent, oily fluid, of sp. g. 1.0929 at 46°, boiling at 370°, miscible with hol and ether, and having a peculiar aromatic smell. In a state of puit may be kept many days under water, in which it is very sparingly ble without decomposition; but when it contains but a minute proportion ee acid or alcohol, it is speedily decomposed into oxalic acid, which is sited in large four-sided prisms, and alcohol. The same reaction ensues an excess of fixed alkali.

#### of Carbonic Ether, or Carbonated Oxide of Ethyl, C4 H5O + CO3.

- ssium being added to oxalic ether, duly warmed, as long as any gas is ved and any excess of the metal removed, the resulting mass was subd to distillation. Carbonic ether was generated, and being conveyed the receiver, formed a superstratum upon the other products of the probeing separated, and freed from water by the chloride of calcium, as redistilled from potassium till, on contact with caustic potash, no oxeculd be formed.
- 616. Carbonic ether is colourless, ethereal, and very liquid, having an ent taste, and an aromatic odour, resembling the ether from which it inates. It is lighter than water, has the specific gravity 0.975 at 66°, at 260°, and burns feebly with a blue flame. It may be mingled in proportions with alcohol and ether, but is insoluble in water. When ed with an alcoholic solution either of the hydrate of potash or soda, it nickly resolved into alcohol and an alkaline carbonate, which separates rater as an oily concentrated solution, or forms as a crystalline powder, water be present. The formation of carbonic ether, which is attended the production of several substances not yet examined, is still unexned.

## Formiated Oxide of Ethyl, or Formic Ether, C4 H5O + C9H O3.

617. To prepare formic ether, seven parts dry formiate of soda are disd with ten parts of sulphuric acid, and six of alcohol, of about 830°.
618. Formic ether is generated, and comes over for the most part withthe application of heat. It is depurated of acid by milk of lime, and

from water by chloride of calcium, which should be added so long as a to

comes moist (5281).

being of the density of 0.912. It boils at 128. Its taste is cooling at spicy. It requires for its solution ten parts of water, but dissolves and proportions in alcohol and ether, in pyroxylic spirit and several fixed selvolatile oils. It is acidified by exposure to air.

#### Of Benzoated Oxide of Ethyl, or Benzoic Ether, C24 H3 O + C4 H4Q.

Liebig, is generated by distilling a mixture of four parts of alcohol, of two parts of crystalized benzoic acid, and one of concentrated, liquid, convolved acid. As soon as the product renders water turbid, the reconstructed benzoic etc. The other, thus obtained, must be added to water to separate it, and be a terwards boiled, with water and litharge, to remove free benzoic acid. In lastly, digested with chloride of calcium.

5621. Benzoic ether is colourless, neutral, and very liquid, having it ethereal but sufficienting odour, and provoking tears. Its specific gravitation, is 1.0539. It boils at 410°, is soluble in alcohol and ether, but a soluble in water. It is decomposed by chlorine, according to Malagon.

producing, among other products, chloride of benzule.

# Of the Tartrate and Citrate of the Oxide of Ethyl, and other " Sala" of Ethyl, so called, of minor importance.

.5622. There are few exacids which may not be united with the exist of ethyl and other exidized compound radicals, so as to form combination a composition analogous to the complex ethers. Those formed with care and tartaric acid, are hardly ethereal in their properties. The extrate quires a heat of 540° for ebullition, and is partially decomposed damages tillation. The tartrate, not being capable of neutralization, is more than gener of sulphovinic acid, viewed as a bibasic acid (5290), than of the complex ethers (3079).

5623. I shall forbear to treat of various compounds, analogous in composition to those above described, whether having the oxide of ethyl, or any other oxidized compound radical, as a base, for reasons like those above

given in relation to the acids (5397).

#### Of Enanthated Oxide of Ethyl, or Enanthic Ether, C. H.O + C. H.O.

to confound it with the oil of wine, Liebig's sulphate of ethyl and etherally distinguished from dilute alcohol. It forms a portion of the constitute about one part in 40,000 of wine. The bouquet by which was wines are distinguished, ought to be ascribed to this etherall compound.

wine, almost intoxicating when plentifully inspired, and a strong disagrable taste. It is soluble in ether and in alcohol, whether concentrated dilute; but not in water. Its density is 362; its volatility feeble. It a quires a temperature between 434 and 446 for obulition. This exist instantly decomposed by fixed caustic alkalies; but not by ammons.

carbonates. When distilled with caustic potash, it is resolved into which comes over, and a very soluble cenanthate of potash.—Gralements.

ple Ethers, formed by the Substitution of another Basacigen Body xygen in the Oxide of Ethyl; or for the Hydrogen in the Water with that Oxide.

Chloride of ethyl, C4 H5 Cl, also called chlorohydric ether, foruriatic ether.—It must be evident, from the comparison of the forove given, with that of ether, C4 H5 O, that this chloride differs t oxide, only in the substitution of an atom of chlorine for an atom :n.

Chloride of ethyl is generated by the distillatory reaction of chloacid, or various chlorides, either with the oxide of ethyl, with alr any other of the compounds of that oxide, such as acetic, citric, artaric ether, &c. Agreeably to one process, alcohol is to be first i with chlorohydric acid gas; according to another, it should be with an equivalent proportion of a strong aqueous solution of that means of a glass retort, communicating by a tube with some water, perature of about 90° F., in a bottle with three orifices. Through æ, the tube proceeding from the retort enters, and is luted air-tight; ther orifice, a tube of safety is inserted; from the third orifice proother tube, arranged so as to communicate, through a refrigerating us, with the interior of a phial surrounded by a freezing mixture. er in the intermediate vessel detains any alcohol or acid evolved ether, which, in consequence of its greater volatility, reaches the The product is freed from water and alcohol by digestion, for twenyours, on chloride of calcium, cooled by ice-water.

Chloride of ethyl is a colourless, ethereal liquid, with an aromatic, ing, and somewhat alliaceous odour. Its density is .874 at 41°. it 52°; does not redden litmus; dissolves in twenty-four parts of roducing a solution which has a fresh, aromatic taste. With solusilver it gives no precipitate. It burns with a bright flame, green order, evolving fumes of chlorohydric acid. In passing through an cent tube, it is resolved into equal volumes of that acid, and olefiant he exposure of this chloride to gaseous chlorine, aided by the solar s given rise to a series of compounds. It is decomposed after some the alkaline hydrates, into a chloride of the ingredient metals,

hol.

Bromide of ethyl, C4 H5 Br, was discovered by Serullas, by dismixture of one part of bromine, four of alcohol, and one-eighth phorus. It is a colourless liquid, denser than water, and very vo-

. Iodide of ethyl, C4 H5 I, may be obtained by distilling alcohol, d with hydriodic acid gas. It is a colourless liquid, of the density 16. It boils at 161°.

. Sulphide of ethyl, C4 H5 S, is formed by transmitting the vapour ohydric ether, through an alcoholic solution of the proto-sulphuret sium; the chlorine being precipitated with the potassium, the sulites with the ethyl and is dissolved, or distils over, if kept suffiwarm. It is a colourless liquid; the boiling point, 185°; density, 1 68°.

# HS.—It might be advantageously called sulphalcohol, since sulphalcohol proper, sulphydre sallocupying the place of water, as may be perceived by the name and in mula.

5683. The best way to prepare this compound, discovered by Zene, is to distil a solution of the sulphovinate of time, of the density of 1.28. In a solution of sulphydrate of the sulphide of potassium, generated by attenting a solution of potash, also of the density of 1.28, with sulphysic and

gus.

5634. The product may be condensed by means of a refrigerator paratus, like that mentioned as having been used for ether (5567). It may be purified from an excess of sulphydric acid, alcohol and water, 50 second distillation from a small quantity of red oxide of mercury, and sesequent digestion with chloride of calcium. Mercaptan bods at 100" see ly, being a colourless, ethereal liquid, with a most penetraring and may portable allianeous odour. Its density is said to be 0.835 at 70°, and 0.842 at 59°. It is soluble in alcohol and ether, but it is very alights ... luble in water. The sulphydric acid of mercaptan reacts powerfolts will metallic oxides, generating water, and a sulphide of the metal. The phide remains in combination with the sulphide of ethyl, thus former class of sulphur salts. The oxide of mercury is instantly converted by mercaptan, into a compound of this class, the mercaptide of mercury, CIP S + Hg S, which is a white, crystalline mass, soft to the touch, want odour, insoluble in water, and fusible, like wax, at 185°. This mercant when distilled leaves cinnabar, and affords a volatile liquid, which has at been examined. The oxide of gold is also strongly acted on by mercaput but other metallic oxides are less affected in proportion as they are Thus, the hydrates of potash and soda have so small electro-positive. reaction with mercaptan. When gently heated, with nitric acid, mercaptan is converted into a new acid, which contains sulphide of ethyl, and it is

ments of sulphuric acid, C<sup>4</sup> H<sup>4</sup> S<sup>6</sup> O<sup>8</sup> (Lœwig, Kopp).

5635. Bisulphide of ethyl, C<sup>4</sup> H<sup>3</sup> S<sup>6</sup>.—A transparent, obvious, which boils at 123.5°, is obtained by distilling a mixture of sulphurents of potash and the persulphide of potassium. It is decomposed by

caustic potash, or by nitric acid (Zeise, Pyrame, Morin).

5636. Selenide of ethyl is obtained, according to Læwig, in the way as the sulphide, substituting in the process, selenide of potassium.

5637. Telluride of ethyl, C4 H5, a very volatile liquid, of a deep control colour (Whoeler), may be obtained also by a similar process, using the

luride of potassium.

5636. Cyanide of ethyl, improperly called cyanhydric ether, — C PN is a colourless liquid, with an insupportable odour of garlic, was classed by Pelouze, by exposing a dry mixture of sulphate of ethyl and rouse is a gentle heat, gradually increased. It has also been formed by deciling in mixture of sulphocyanide of potassium, alcohol, and sulphuric acid. It at thick, oily liquid, of the density of 0.7, boiling at 179.6°.

Of the Dehydrogenation and Oxidation of Ethyl, as contained in Ether or Alcohol, and of the Oxidation of the residual Products.

5639. The precipitation of carbon which gives a fuligiaous character to the flame of essential oils and resins,
has been ascribed to an inadequate supply of oxygen, and
the superior affinity of hydrogen for oxygen, comparatively with carbon, at moderate temperatures. In consequence
of this preference thus given, when some of the compounds
of ethyl are subjected to oxidating agents, in processes
below the temperature of ignition, more or less hydrogen
is removed according to the intensity of the reaction.
Thus, from alcohol C<sup>4</sup> H<sup>5</sup>O + HO, two atoms of hydrogen
being taken, aldehyde is engendered, C<sup>4</sup> H<sup>3</sup>O + HO. These
elements, by the absorption of one atom of oxygen, form
an acid which has been called aldehydic acid, or acetous
acid, C<sup>4</sup> H<sup>3</sup> O<sup>3</sup> + HO. Another atom of oxygen converts
this acid into acetic acid, C<sup>4</sup> H<sup>3</sup> + O<sup>3</sup> + HO. Aldehydic
acid has also been designated as acetous acid, having the
same radical, and less oxygen than acetic acid (3094).

5640. Acetyl, of which the formula is C<sup>4</sup> H<sup>3</sup>O, has been already described as a compound radical, indebted for its existence to the removal of two of the five atoms of hydrogen belonging to ethyl (3093). Liebig attributes this result to the oxidation of the ethyl; but it is, as I conceive, a case of dehydrogenation of ethyl, resulting from the oxidation of two-fifths of its hydrogen. The relation between this radical and its progeny may be seen in the following

table.

Acetyl, - - - C<sup>4</sup> H<sup>3</sup>

Aldehydic, or hydrate of the oxide of acetyl, - - - C<sup>4</sup> H<sup>3</sup>O + HO.

Acetous, or aldehydic acid, - C<sup>4</sup> H<sup>3</sup>O<sup>2</sup> + HO.

Acetic acid, - - - C<sup>4</sup> H<sup>3</sup>O<sup>3</sup> + HO.

No oxide of acetyl has been ascertained to exist uncombined with water and corresponding to common ether.

Of the Hydrated Oxide of Acetyl, called Aldehyde.

5641. By inspection of the respective formulæ, it will be perceived, that this compound differs from alcohol only

in the loss of two atoms of hydrogen. Hence its name from the first syllable of each of the words alcohol and dehydrogenatum. Aldehyde is one of the products of the decomposition of alcohol, or ether, by passage through a tube at a low red heat: during etherification by nitric and (5586): by platina wire in the lamp without flame, and other cases. Liebig's process for the preparation of allehyde is as follows:—Six parts of oil of vitriol with four of water; four parts of spirits of wine and six of black and of manganese, are to be distilled with a very gentle beat and the product collected in a receiver surrounded will ice-water. This process is completed as soon as the terials in the retort cease to froth up. Kane observes that a purer product may be obtained by distilling two parts of spirits of wine with three of bichromate of potash, three of oil of vitriol, and six of water; the two last being previous ly mixed and allowed to cool. To obtain aldehyde absolutely pure, it must be combined with ammonia; the resuling crystallized ammoniacal compound must afterward be decomposed by dilute sulphuric acid, distilled, by means of a water-bath at 120°, with great care, and finally recubel over fused chloride of calcium.

5642. Aldehyde is a colourless, inflammable liquid, of a peculiar ethereal and suffocating odour. It boils at 72, has the density 0.790 at 64.40, and is soluble in water, alcohol and ether. By absorbing atmospheric oxyges, it is converted spontaneously into acetic acid. It dissolves phosphorus, sulphur and iodine. Aldehyde is capable of combining directly with ammonia and potash, thus evencing

an approach to the acid character.

5643. Ammoniated Aldehyde, or the Hypoacetite of Ammonia, C<sup>4</sup>H<sup>3</sup>O, NH<sup>3</sup> + HO. In this compound aldehyde appears to act as an acid in entering into union with the cure of ammonium (1106), so that it might be consistently designated as hypoacetous acid. Ammoniated aldehyde appears in acute, colourless, transparent, brilliant, friable rhomboids, of a considerable magnitude, and which have an odour of spirit of turpentine. These crystals fuse between 150° and 172°, and distil without decomposition. If they burn with a yellow flame. In the air, or even in closed phials, they turn brown, acquiring the smell of a burnt feather. Under pure ether they may be preserved, but not for a long time. These crystals are soluble as

water in all proportions, and more readily in hot than in cold alcohol. In ether they are but little soluble.

5644. Acetal, a compound of aldehyde with ether, C<sup>4</sup> H<sup>5</sup>O + C<sup>6</sup> H<sup>6</sup>O + HO = C<sup>8</sup> H<sup>8</sup> O<sup>5</sup>, was discovered by Dobereiner, and described as oxygenated ether. It is formed by the reaction of platinum black with the valour of alcohol, with the presence of oxygen. Acetal is a colourless liquid, taving a peculiar odour, suggesting that of Hungary wines. It boils at 103°; its density is 0.823 at 68°. It is soluble in six or seven parts of water, and mixes with alcohol in all proportions.

5645. The crude formula of acetal being C<sup>8</sup> H<sup>9</sup> O<sup>8</sup>, two atoms of it will be found to contain the ingredients of three atoms of ether, and one

of acetic acid.

5646. Resin of aldehyde is a product of the decomposition of aldehyde

by alkalies, with the assistance of air.

5647. Elaldehyde.—When pure anhydrous aldehyde is kept for some time at 32°, while gradually losing its power to mix with water, it is transformed into a coherent mass of long, transparent, needle-shaped crystals, resembling spiculæ of ice. This is elaldehyde, which is similar in composition to aldehyde, but of three times the atomic weight, judging from the lensity of its vapour. Elaldehyde fuses at 35.6°, and boils at 201.2°.

5648. Metaldehyde is another product of the condensation of the elements of aldehyde, which appears in aldehyde left for some time in a well-stopped phial, in the form of white and transparent needles, or colourless prisms, which gradually attain a certain magnitude. It sublimes at 248°, without fusing, and condenses in the air in snowy and very light flocks. It is insoluble in water, but dissolves easily in alcohol.

Of some interesting Results of the substitution of Chlorine, Bromine, Sulphur, and other Basacigen Bodies, for the Hydrogen or the Oxygen in the Compounds of Ethyl and Acetyl.

5649. Of the Chlorohydrate of the Chloride of Acetyl, Chlorine Ether, Bicklorine Ether, C<sup>4</sup> H<sup>3</sup> Cl + Cl H.—Under the head "olefiant gas" [1274], it was mentioned (1292) that olefiant gas received its name from its capacity of forming a liquid of an oily consistency, having an agreeable smell and taste. On account of its resemblance to ethereal compounds, as respects fragrance, solubility and taste, and the presence in it of two atoms of chlorine, it has been called bicklorine ether. Considering olefiant gas a hydruret of acetyl, C<sup>4</sup> H<sup>3</sup> + H, the liquid in question is, by Liebig, reated of under the appellation at the head of this article, which indicates it to consist of chlorohydric acid, C H, and chloride of acetyl, C<sup>4</sup> H<sup>3</sup> Cl.

of moist chlorine and the gas above mentioned, within a large receiver, over water. Liebig recommends the reaction of the same olefant gas with the perchloride of antimony, so long as there is any absorption. The resulting mass is to be subjected to distillation, till the product ceases to yield any ethereal liquid on the addition of water. The combination thus elaborated, requires to be depurated by redistillation with water, and subsequent agitation with sulphuric acid to depurate it of moisture. This ordeal is to be repeated until it ceases to be affected by sulphydric acid, or to emit chlorohydric acid during distillation. Finally, being successively washed with water, and kept in contact with chloride of calcium, it becomes quite pure.

5651. Thus obtained, the chlorohydrate of the chloride of acety are lourless, very liquid, and, as respects smell and taste, sweet and agree It boils at 179°. It may be distilled, without decomposition, from the the line hydrates. It communicates its smell to water, although matter therein. In alcohol and ether it dissolves in all proportions. By an are holic solution of potash, it is decomposed into chloride of potash and cheride of acetyl. It was by the exposure of this chlorohydrate to recommitte the chloride, in the sunshine, that Faraday (1242) obtained the exposure of carbon.

5652. Chloride of Acetyl.—This is a gaseous product, of which the tion is above made as resulting from the reaction of its chlorohydrate who potash, in alcohol. It has an alliaceous odoor, and may be injuried at the

temperature of -6%.

5653. Bromohydrate of Bromide of Acetyl, Bromide of Acetyl, his hydrate of lodide of Acetyl.—Substances are described by Lucky, to with these names are given, which indicate their analogy with the two compounds of chlorine last described.

5654. Chloroplatinate of Chloride of Acetyl.—By this name a compound has been designated, in which chloroplatinic acid takes the place of

chlorohydric acid in the chlorohydrate of the chloride of acetyl.

to a colourless oily tiquid, which results from the saturation of activities ether with chlorine, desiccated by being passed through concentrated phuric acid. The formula of ether being C\*H\*O, two out of the five second from supply their place. Thus the oxychloride of acety I is generated process analogous to that by which acetic acid is generated, oxygen forming, in one case, the same part as chlorine in the other.

above given, the ingredients of acetyl combined both with oxygen and the rine. It is therefore designated as an oxychloride, although uses the name has been appared to the union of an oxide and a chloride, each see-

rally combined with distinct atoms of the same radical.

occupies the place which chlorine fills in the oxychloride of acetyl, and of course that which oxygen fills in anhydrous acetors acid. The substants of sulphur is effected by the reaction of the exychloride of acetyl with an phydric acid; during which, the two atoms of chlorine uniting with the coordinate of hydrogen, two atoms of sulphur supply the place of the chlorine this is moved. During this last mentioned reach in another compound is feasien which only one atom of the chlorine in the exychloride is replacely sulphur. The formula, of course, is C\* H\* C\* SO. The cityl mastic other may, by reaction with charme have its hydrogen so far replacely chlorine as to be converted into an oxycle made, while its union with acetic acid remains unbroken. Benzoic other may not only have as the similarly changed, but the oxygen, forming its acid with benzule, may be placed by chlorine; so that for Bz O + C\* H\* O, a compound results, to presented by Bz Cl + C\* H\* Cl\* O.

5658. Chloroxalic Ether, C\* Cl\* O + C\* O\*.—This ether is created by the substitution of chlorine for the whole of the hydrogen in oxalic ether, of which the formula is C\* H\* O + C\* O\*. It will be seen, on comparing thereformulæ, that they differ only in this; five atoms of chlorine in one are stituted for a like number of hydrogen in the other. It was obtained by

jecting oxalic ether in a retort, surrounded by boiling water and exposed

the solar light, to a current of chlorine.

5659. Chloral,\* C4 H3O + HO, is the name given to a compound disrered by Liebig, in which all the hydrogen of aldehyde is replaced by orine. It might, with propriety, I think, be called hydrated oxide of loracetyl, chloracetyl being understood to apply to the compound C4 Cl3, ich takes the place of acetyl as the organic radical. Chloral is the imate product of the dehydrogenation of anhydrous alcohol, by dry lorine, and the substitution of three atoms of chlorine for five of hy-

gen.

5660. By subjecting, for twelve or fifteen hours, anhydrous alcohol to lorine, dried by passage through sulphuric acid, a copious evolution of lorohydric acid ensues, and a dense oily liquid is generated, which conals on cooling, being an impure hydrate of chloral. It is requisite to eck the reaction in the first instance by immersion in water, afterward to sist by heat the expulsion of the chlorohydric acid. The hydrate is puied first by heating it in a well-stopped flask, with nearly three times its Ik of sulphuric acid, when the chloral, depurated of water, forms a supertant stratum. This being separated, and boiled to expel free chlorohyic acid, or alcohol, to remove any residual moisture, the chloral should distilled from an equal volume of sulphuric acid. Finally, it must be stified from lime, which, after being slaked, was rendered anhydrous by posure to a bright red heat.

5661. Chloral, thus obtained, is a dense, oily, colourless liquid, greasy the touch, having a penetrating, disagreeable odour, which provokes Its taste is at first greasy, then caustic. It produces on paper an anescent stain. Its density, at 64.4° is 1502; its boiling point is 201°, arly. It may be distilled without alteration. Its vapour is nearly five nes as dense as that of air, its equivalent being four volumes. Chloral is iscible either with alcohol or ether. Aided by heat it dissolves sulphur,

osphorus, or iodine, apparently without alteration.

#### OF METHYL ETHERS.

### Of the Oxide of Methyl, or Methylic Ether, C<sup>2</sup> H<sup>3</sup> O.

5662. In treating of the hypothetical compound radical, nethyl, it was stated, that it was considered as performing, 1 the compound above mentioned, a part analogous to

nat which ethyl is inferred to perform in alcohol.

5663. The oxide of methyl is prepared by distilling one art of wood spirit, with four parts of sulphuric acid, the esulting gas being transmitted, successively, through a ottle containing milk of lime, and several bottles duly harged with pure water. In this liquid the gas dissolves, nd being evolved by a boiling heat, may be collected over nercury. Oxide of methyl is an inflammable, colourless

<sup>\*</sup> The account of chloral, given under the head of inorganic compounds of carbon. sing very brief, I have not hesitated to treat of it again, as an organic product, itheat reference to that imperfect notice.

gas, of an agreeable ethereal odour. For liquefaction it requires a temperature below 3.2°. Thirty-seven volume at a of this gas dissolve in one of water. Alcohol, hydratedor ide of methyl, and concentrated sulphuric acid, take ne to a greater extent. From the latter it separates on divtion with water. The density of the gas is, by expenses, 1605; by calculation, 1570; its combining measure being two volumes.

5664. By combining with the vapour of anhydron \* phuric acid, in a glass balloon, carefully cooled, the oxid of methyl forms a neutral sulphate. (Regnault.)

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100

Of Hydrated Oxide of Methyl, called Pyroxylic, or Want Spirit, Methylic Alcohol, C2 H3 O + HO.

5665. In the process of purifying acetic acid from wood (5199), the crude acid is saturated by lime, and concentrated by distillation, of which the first product contains the cmiss wood spirit, which may be partially purified by record distillation from quick-lime; and is found in this state # commerce. It is still a heterogeneous mixture, contained besides the hydrated oxide of methyl, which forms the larger part of it, acetone, and several other volatile and inflammable liquids. To purify the spirit in question, it treated with an excess of chloride of calcium, in a retor, and distilled by a water-bath heat, which expels the more volatile liquids, and leaves the wood spirit in union with the chloride of calcium. A volume of water, equal to the volume of wood spirit employed, is then added to the retort, and the distillation continued. The spirit comes over imbued with a small quantity of water, from which # may be completely separated by subsequent distillance from quick-lime (5711, &c.).

5666. Wood spirit is a volatile, colourless liquid, sindtaneously recalling the taste and odour of acetic ether w alcohol. It is very inflammable, and burns with a pale flame. It mixes with pure water without becoming turbed, and likewise with alcohol and other. Its density is 0,798 at 68°; its boiling point, 140°. The density of its vapour is by experiment, 1120; by calculation, 1100; its combining

measure or equivalent comprises four volumes.

5667. With the aid of heat, hydrated oxide of methyl dissolves small portions of sulphur and phosphorus, and may also serve as a solvent for the resins employed

paking varnishes. It mixes with volatile oils. Wood spirit is, like alcohol, acted upon by chlorine, peroxide of manganese or sulphuric acid, and by oxidizing agents in general, and yields analogous products. It is also decomposed by potassium, with disengagement of pure hydrogen.

5668. Anhydrous barytes, although not soluble in alcobol, dissolves in pure wood spirit, with much heat, and forms a compound, C<sup>2</sup> H<sup>3</sup> O HO + Ba O, which crystallizes in needles of a silky lustre. Lime is likewise soluble

in wood spirit.

5669. Chloride of calcium dissolves eagerly in this solvent, so as to cause much heat. From a concentrated polution, it crystallizes in large, deliquescent, hexagonal tables, which contain two atoms of hydrated oxide of methyl, united with one atom of chloride of calcium.

5670. Neutral Sulphated Oxide of Methyl, C' H' SO'.—This member of the methyl series, which has no analogous compound in that of ethyl, is generated either when oxide of methyl and anhydrous sulphuric acid are brought into contact, or when one part of the hydrated oxide is distilled with eight or ten parts of sulphuric acid; the product being purified by washing with water, and distillation from chloride of calcium and quicklime, successively. Sulphated oxide of methyl is a colourless liquid, of an alliaceous odour, of density 1.324 at 71°.6. It boils at 370°.4, and may be distilled without change. The density of its vapour is 4363.4; it consists of equal volumes of anhydrous sulphuric acid and oxide of methyl, condensed into one volume; its combining measure being four volumes, the same as that of oxide of methyl. It is very slowly decomposed by water when cold, but rapidly when hot; the acid sulphated oxide of methyl and water being formed, while oxide of methyl is liberated. By double decomposition this compound may be employed in preparing all the other compounds of methyl.

5671. Acid Sulphated Oxide of Methyl, Bisulphated Oxide of Methyl, Sulphomethylic Acid, C<sup>2</sup> H<sup>2</sup>O + S<sup>2</sup>O<sup>6</sup> HO. This congener of sulphovinic acid, discovered by MM. Dumas and Peligot, and by Dr. Kane, about the same time, is formed by mixing concentrated sulphuric acid with hydrate of oxide of methyl; or by dissolving the neutral sulphate in boiling water. Obtained by the latter method, and concentrated by evaporation, it is a colourless, syrupy, and very acid liquid; which, in dry air, becomes a mass of white crystalline needles. It combines with bases forming double salts, in which the basic water of the acid is replaced by a metallic oxide.

These double salts are soluble in water.

5672. Nitrated Oxide of Methyl, Me O NO5.—To prepare this compound, one part of nitrate of potash, and a mixture of two parts of concentrated sulphuric acid with one part of wood spirit, are introduced into a retort. The mass rises in temperature, and a liquid distils without additional heat. For its condensation, a refrigerated tube should be employed, terminating in a refrigerated flask. The heavier of the two liquids found in the flask is nitrated oxide of methyl, contaminated by a portion of a very volatile compound, supposed to be formisted oxide of methyl, which imparts the

odour of cyanhydric acid. The product is rectified from chlorde of a cium and from litharge. The last portions which distil over are perceipture. Nitrated oxide of methyl is a colourless liquid, of a weak, condour, which burns readily with a yellow flame; its density is 1.-22 at 71°.6, and boiling point 150°.8. Heated above 246°, its vapour is desposed explosively, producing carbonic acid, water, and deutoxide of an all proportion with alcohol, ether, and wood spirit.

### Of the Hyponitrite of the Oxide of Methyl, or Methylic Hyponium Ether, "Nitrite d'Oxide de Methyl" of Liebig, and others.

5673. In his late Treatise on Organic Chemistry, Liebug makes the following statement:—"The reaction which nitric acid excremes with the hydrated oxide of methyl, is not like that which it excremes with the hydrated oxide of methyl, is decomposed with great difficulty, giving beauties, while this liquid is decomposed with great difficulty, giving beauties oxide of methyl is not altered by nitric acid, unless at a toking beauties of methyl is not altered by nitric acid, unless at a toking beauties and generated, but no hyponitrite ("nitrite") nor nitrate of the oxide of methyl has not exist." Traite, 552.

8674. Last winter, however, Dec. 1841, I found that by subjecting put wood spirit to the process already described for producing the hyportest ethyl, a congenerous ethereal product was obtained (55%3). Hyperitest methyl has a great resemblance to its congener above named, in own, amell, and taste; though there is still a diversity sufficient to emphis a constant.

ful observer to distinguish one from the other-

troducing the refrigerated materials into a bottle surrounded be as and water, was resorted to, substituting wood spirit for alcohol, it was less that the other did not separate from the spirit in completely as in the passes a which alcohol was the material. This I ascribe to the atheir instance water and wood spirit being inferior to that between this last instance quid and alcohol. The boiling point of both of the others seemed to be nearly the same, and in both, in consequence of the escape of an ethered gas an effervescence resembling that of could too, was observed to take passes at a lower temperature, than that at which the bounts point became a bonary.

efforts to produce the methylic hypomtrous other had facted. The inhom of others, and my success, cannot excite suprise, when the differenced be habitudes of wood spirit and alcohol, with intric acid and alcohol, or accident to view, and the difference between my process and those followed a farrope, by which more or less natic acid is brought into contact with the prit employed. When alcohol is presented to notice acid, a reciperation composition ensues. The acid loses two utoms of oxygen, which, by taking two atoms of hydrogen from a portion of the a cohol, transforms it into be dehyde, while the hypomtrous acid resulting mevitably from the parts of oxidizement of the natic acid, unites with the base of the remaining part of the alcohol. But when pyroxylic spirit is presented to intric acid, the act without decomposition, combines with methyli, the base of thus hydrer bence, as no hypomitrous acid is evolved, no hypomatric can be predicated.

Thus in the case of the one there can be no ethereal hyponitrite, in that of the other no ethereal nitrate.

5677. Oxalated oxide of methyl, C<sup>3</sup> H<sup>3</sup> O, C<sup>3</sup> O<sup>3</sup>, is a white, transparent, and brilliant mass, composed of rhomboidal tables, which fuses at 123°.8, and boils about 321°.8. It is decomposed by water, and resolved into hydrated oxalic acid and wood spirit.

that for the formiated oxide of methyl is obtained by a process analogous to that for the formiated oxide of ethyl (5617), substituting pyroxylic spirit for alcohol. It is lighter than water, and boils between 96 and 100°; its odour

suggesting that of acetic ether.

\* Reaction of Chlorine, Iodine, Cyanogen, and Sulphur, with Methyl and its Compounds.

reaction of chlorohydric acid with hydrated oxide of methyl: the rationale being the same as when this acid is presented to a hydrated metallic protoxide. But it is best obtained, as are all the halogen compounds of methyl, by distilling the haloid salt, containing the halogen body with which the union is to be made, with a mixture of sulphuric acid and hydrated oxide of methyl: of course, in the case in point, chloride of sodium may be used. Chloride of methyl is a colourless gas, of an ethereal odour and sweet taste, having the density 1737.8 by experiment, and 1729 by calculation; the combining measure being four volumes. Water dissolves 2.8 volumes of this gas at 60°.8. It is not liquefied by a cold of —0.4. It should be remembered that the chloride and oxide of methyl, are both much, more volatile than the chloride and oxide of ethyl (873).

5680. Iodide of Methyl, C' H'I.—This is a colourless liquid, which inflames with difficulty, and boils between 104° and 122°. Its density is

2.337 at 69.8.

5681. Fluoride of Methyl, C<sup>a</sup> H<sup>a</sup> F, is obtained by distilling sulphated oxide of methyl with fluoride of potassium. It is a colourless gas, of which the density is 1186; and, for the solution of which, fifteen volumes of water are requisite.

5682. Cyanide of Methyl, C' H' Cy, is an ethereal liquid, insoluble in

water.

5683. Sulphide of Methyl, C<sup>2</sup> H<sup>2</sup> S, is a very limpid liquid, of which the odour is extremely disagreeable. Its density is 0.845 at 69°.8, and its boiling point 105°.8. The density of its vapour is by experiment 2115, by theory 2158; its combining measure being two volumes. Sulphide of methyl is formed by double decomposition, when chloride of methyl is transmitted through an alcoholic solution of protosulphide of potassium.

5684. Sulphydrate of the Sulphide of Methyl, or Methylic Mercaptan, is a colourless liquid, lighter than water, which boils at 69°.8 and acts on

oxides of mercury and lead like sulphydrate of sulphide of ethyl.

5685. Compounds having methyl for their radical, correspond so closely with those in which ethyl sustains the same character, that knowing the history of one class, it is easy to imagine the properties of the other. Anhydrous metallic salts do not alter them, while the hydrated alkalies disenges hydrated oxide of methyl from them with great facility.

5686. Chlorine decomposes the gaseous oxide of methyl, forming chlorohydric acid, and the following products, as observed by M. Regnault:—

Monochlorinated oxide of methyl, - C<sup>2</sup> H<sup>2</sup> Cl O

Bichlorizated oxide of methyl, - - C\* H Ct\* O
Perchlorizated oxide of methyl, - C\* Ct\* O

5687. Chlorine is absorbed with great avidity by hydrated oxide of the thyl, a heavy oil being generated, which has not been well examend.

5688. The reaction of chlorine with chloride of methyl, is the same describes of compounds, in which, generally, the proportion of chiefes the creases as the reaction is prolonged.

Chloride of methyl, - - - C\* H\* C

Monochlorinated chloride of methyl, - C\* H\* C

Bichlorinated do. (chloroform), C\* H\* C\*

Perchlorinated do. - - C\* C\* C\*

5689. The monochlorinated chloride of methyl has an odour which a very sharp, but is, in other respects, similar to the oil of olefant get. Its tilled with an alcoholic solution of potash, a trifling precapitate of chlorad potassium is formed, and it comes over unchanged.

5690. The perchloride of carbon, Co Clo, which is named, above, perchlorizated chloride of methyl, is not altered by a solution of sulphydrate of potassium. It is decomposed by heat, yielding different chlorides of second

bon according to the temperature.

5691. At a low red heat, this chloride, C\* Cl\*, appears to be committed into another chloride of carbon, C\* Cl\*, supposing the combining manner of the latter to be four volumes, its density being 4092. This new darks of carbon must therefore be isomeric with Faraday's sesquichloride, C\* U\*, but of only half the density. When decomposed at a higher temperature, it gives small silky crystals, constituting the chloride of carbon, C\* C\*. Ch. Lastly, at a bright red heat, the liquid chloride of carbon, C\* C\*. at the product.

5692. Chlorine acts readily upon the sulphide of methyl, and open to compounds of the oxide of methyl with acids, constituting the compound methylic ethers. A benzoate and acetate of an oxychloride of forest have been produced, having the following formulæ:—

 $C^{\alpha}HC^{\alpha}O+BzO$ .  $C^{\alpha}HC^{\alpha}O+AcO$ .

From a mixture of iodine, nitric acid, and wood spirit, left to isself for a long time, yellow crystals are deposited. Bromine, under the same estances, yields a heavy only liquid.

### Of Formyl Ethers.

5693. Hydrated oxide of methyl, when brought into contact with plant mum black and atmospheric air, is converted into pure formic acid, is the substitution of two atoms of oxygen for two of hydrogen. The chart effected is, therefore, perfectly similar to that by which alcohol is, is the name agent converted into acetic acid. Oxide of methyl, formula C HU + HO and 40, is equivalent to formic acid, C° HO + 3HO. Hence the inference, that formic acid contains a radical formyl, C°H, to which a the the same relation as acetic acid has to acetyl: acetic acid, C° H° + O° formic acid, C°H + O° (4019).

5694. Formyl is the hypothetical radical of the following compounds-

# Of Methylal, C<sup>6</sup> H<sup>8</sup> O<sup>4</sup>, a Compound of Hydrated Oxide of Formyl, with Oxide of Methyl.

5695. By distilling two parts of wood spirit with two parts of peroxide of manganese, and three parts of sulphuric acid, diluted with three parts of water, Dr. Kane obtained a substance mixed with several other bodies, which he named formomethylal. It was considered a tribasic formiated mide of methyl, but was afterwards shown by Malaguti to be a mixture of brimiated oxide of methyl and a particular substance which he named methylal. To purify the methylal from the formiated oxide of methyl, the

atter must be decomposed entirely by hydrate of potash.

matic odour; which is miscible with three parts of water, and may be separated from that liquid by chloride of calcium, or hydrate of potash. It is rery inflammable, and burns with a white flame. The density of methylal s 0.8551; its boiling point 107°.6; its combining measure contains four volumes. Methylal may be represented as a compound of one atom of hylrated oxide of formyl with two atoms of oxide of methyl = C° HO, HO + 2C° H° O. Regnault has explained its formation, by supposing that hree atoms of oxide of methyl, formed by the action of sulphuric acid upon bydrated oxide of methyl, group together so as to form a single molecule = C° H° O°. This molecule, by exposure to peroxide of manganese, loses one atom of hydrogen, gaining one of oxygen, so that the compound C° H° O° results. The formation of acetal, which corresponds with methylal in the acetyl series, is explained by Regnault in the same manner.

5697. Artificial Oil of Ants, C<sup>5</sup> H<sup>2</sup> O<sup>2</sup> (Stenhouse).—This name was applied, by Dobereiner, to an oil generated during the preparation of formic scid. It was obtained by Dr. Stenhouse in larger quantity than it is proluced during the ordinary process, by distilling a mixture of equal weights of out-meal, or saw-dust, and sulphuric acid diluted with its own bulk of water. In the process for formic acid, the peroxide of manganese cannot be omitted without greatly reducing the product; but in the process in question it should be left out. When oil of ants is purified, the taste and smell are very pungent and aromatic, resembling that of oil of cassia. It burns very readily with a bright yellow flame. Its density is 1.1006 at 80°.6; ts boiling point 334°.4. It is soluble in water, but more so in alcohol and ther. It is decomposed by potassium with effervescence; but neither the

nqueous nor the alcoholic solution of potash is affected by it.

### Compounds of Formyl with Chlorine, Bromine, Iodine, and Sulphur.

5698. Protochloride of Formyl, C<sup>2</sup> H Cl.—One of the substances which Regnault obtained by the reaction of chlorine with chloride of acetyl, name-y, C<sup>2</sup> H<sup>2</sup> Cl<sup>2</sup>, is considered by Liebig as the protochloride of formyl, its tomic weight being divided by two.

5699. Bichloride of Formyl, C<sup>2</sup> H Cl<sup>2</sup>.—According to Liebig, of one of be combinations, generated by the reaction of chlorine with the chloride of thyl, the formula is C<sup>4</sup> H<sup>2</sup> Cl<sup>4</sup>. This being divided by two, gives that of

he bichloride, as above stated.

may be made, by exposing a mixture of chloride of methyl, C<sup>1</sup> H<sup>2</sup> Cl, and chlorine to the direct rays of the sun; by distilling chloral with brown water, or milk of lime, but more conveniently by distilling a dilute material of hypochlorite of lime, or bleaching salt, with acetone, alcohol, or with apirit. For this purpose, one part of slaked lime is suspended in trees four parts of water, and impregnated with chlorine till the greater part of the lime is dissolved. The lime must be in sufficient excess, browner, to render the liquid slightly alkaline. When the solution of hypochlora this made has become clear, \(\frac{1}{34}\) of its volume of alcohol should be with the audiculated to the distillatory process, at a gentle heat, by means of a concious retort. The product, consisting of perchloride of formyl, many alcohol, being agitated with water, the perchloride separates as a decomposition, and may be obtained perfectly pure by digesting it upon chlorae acid.

ethereal odour, and sweetish taste; its density is 1.480 at 64°.4; is how point, 141°.44. It is difficult to inflame, but burns in the flame of a less imparting a green colour. An alcoholic solution of potash converts to formiate of potash, and chloride of potassium, on which the mass chroform is founded. Fo Cl<sup>3</sup> and 4 Po O = Po Fo O<sup>3</sup> and 3 Po Cl. The sity of its vapour is, by experiment, 4200; by calculation, 4116; a cabbining measure is 4 volumes. Chloroform may be distilled from schools acid, potassium, or potash, without being sensibly altered. Export of chlorine to the direct rays of the sun, it is decomposed, and converted at 172°.4, and of which the density of the vapour is 5300, and constant measure four volumes. This chloride results from the substitutes of the rine for the whole of the hydrogen and oxygen in formic acid, C° Cl<sup>2</sup> while the well known sesquichloride of carbon, C<sup>4</sup> Cl<sup>2</sup> + Cl<sup>2</sup>, is smally

derived from acetic acid.

5702. When the above described chloride of carbon is made to pass in vapour through a porcelain tube, at a low red heat, it is resolved into two new chlorides of carbon, of one of which the composition is CCI, while of the other the composition is CCI, according to Regnault.

5703. Chlorohydrate of the chloride of formyl, 2 C H Cl H Cl, is an of the products of the reaction of chlorine, with the chlorohydrate of in

chloride of acetyl.

5704. Perbromide of formyl, bromoform, C<sup>2</sup> H, is prepared like the prechloride, and very analogous to it in properties. Its density is 2.10. Em less volatile than the perchloride, and more easily decomposed by a ballow.

stance discovered by Serullas, which is often described as an iodide of explora. To obtain it, an alcoholic solution of potash is added to a solution of iodine in alcohol till the last is decolorized, carefully avoiding any excess the alkali. The alcohol being allowed to escape by gentle exaporation, the iodide of formyl is deposited in crystals, which are purified from solide of potassium by washing with pure water. This compound results from the reaction of one atom of alcohol, with six atoms of potash and eight areas of iodine, by which one atom of periodide of formyl, one atom of forming of potash, five atoms of iodide of potassium, and four atoms of water, are formed.

| 1 atom of alcohol,                                           | • | • | • | • | C4 | H O H                         | }                              |
|--------------------------------------------------------------|---|---|---|---|----|-------------------------------|--------------------------------|
| 8 atoms of iodine,                                           | • | • | • | • |    |                               | Ia                             |
| 6 atoms of potash,                                           | • | • | • | • |    | O <sub>6</sub>                | Po                             |
|                                                              |   |   |   |   | C• | H <sub>6</sub> O <sub>8</sub> | Is Pos                         |
| 1 atom of periodide of formyl, 1 atom of formiate of potash, |   |   |   | • | •  | C <sub>a</sub> H              | Is.                            |
|                                                              |   |   |   | • | •  | C <sup>a</sup> H              | O <sup>4</sup> Po              |
| 5 atoms of iodide of potassium,                              |   |   | • | • | •  |                               | I <sup>s</sup> Po <sup>s</sup> |
| 4 atoms of water,                                            | • | • | • | • | •  | H• (                          | 04                             |
|                                                              |   |   |   |   | •  | C• He                         | Os Is Pos                      |

5706. Iodoform crystallizes in brilliant yellow plates: has a characteric odour suggesting that of saffron; is insoluble in water, but very soluble alcohol, ether, and wood spirit. It sublimes at 212°, and at 248°, is solved into carbon, iodine, and hydriodic acid. When distilled with loride of phosphorus, or with corrosive sublimate, it yields a peculiar juid, of a deep red colour, and a density of 1.96, which contains chlorine, dine, and formyl.

5707. Sulphide of formyl, C' H' S' (Bouchardat), is a liquid obtained distilling one part of iodide of formyl, with three parts of sulphide of ercury. By reaction with hydrate of potash it may be converted into sul-

side of potassium and formiate of potash.

### Of Xylite, or Lignone.

5708. Having received from Dr. Ure a bottle of a liquid, which I underpod to be pure wood spirit, I subjected it to the usual test of saturating it
ith chloride of calcium, with which wood spirit reacts eagerly, generating
at as already mentioned. I found, however, that a colourless liquid
parated, and formed a supernatant stratum, in which the chloride above
med did not appear to be soluble.

5709. When this liquid, and the solution of the chloride in wood spirit, ere subjected to the distillatory process, by means of a boiling water bath, ily the former come over, the wood spirit being retained by the chloride.

5710. It seems from the account given by Graham, 836, that the liquid hich was distilled, has been examined by Wiedman and Schweiger, and scribed under the names at the head of this section. The formula asymed to it is C<sup>6</sup> H<sup>6</sup> O<sup>2</sup> 1; which, as the admission of half atoms is inconstent with the grounds on which the atomic theory is built, should be subled.

5711. The boiling point of xylit is about 142°, its density 0.816. The maity of its vapour to that of atmospheric air as 2177, by experiment, ad 2159 by calculation. Pure xylit has an agreeable, sharp, empyreuatic taste. It is soluble in water, and burns with a white flame.

5712. Mesiten, C<sup>6</sup> H<sup>5</sup> O<sup>2</sup>, agreeably to the same authority, is a liquid tained by distilling equal parts of xylit and sulphuric acid. Chloride of deium is utterly insoluble in mesiten, of which the boiling point is 145°, and the density 0.808.

5713. Mesite is the name given to a liquid which is a concrete product the destructive distillation of wood, which gives birth to wood spirit and

to xylit. Being less volatile than the last mentioned product, it comes at later, and hence it may be isolated. It is formed also by the reacted

xylite with potash and potassium.

of potash with mente. It is in its properties etherend; being colored very liquid, and having the odour of peppermint. It is but alight; such in water, but very soluble in alcohol, in wood spirit, xylit or ether. It had at 230°, and burns with white smoky flame.

5715. Xylite oil is produced from xylit naphtha, by a renewd and

tion with hydrate of potash. This oil has ethercal properties.

5716. Methal, C' H', is a liquid generated by the reaction between siphuric acid and xylit. Pyroxanthin, by the distillation of crude want of from slaked lime. These substances are more of the nature of an exist oil, or camphor, than of that of an ether.

### Of the Ethereal Compounds of Mesityl, or Mesitylene.

5717. The origin and characteristic properties of mesity! were set in the general account of it, as one of the compound radicals among well it stands distinguished as being one of the few which are capable of set tion. As respects its properties, it may be considered as an other, you

5718. Of the Chloride of Merityl, Co Ho Ch.—I give precedence as a compound over the exide, contrary to the course pursued in the otherwise as it is only by means of the former that the latter has been character are mixed gradually with one of acctone in a refingerated vessel. On a addition of water to the resulting mass, an oily liquid separates, which a sufficiently heavy to sink in water, and which heat resolves into characteristic acid and mesityl. This oily liquid is the chloride of the last mention radical.

the alcoholic solution of the chloride, above described, with cause made in excess; followed by the addition of a large quantity of water, and liquid separates, which, being desiccated by contact with chloride of recium, is afterwards distilled. Thus purified, it is a colourless found, true the odour of peppermint. It bods at 24%, is inflammable, and burns and brilliant flame, attended with much smoke.

a mixture of acctone, phosphorus, and todine, to the distillatory process

5721. Chloride of Pteleyle.—When mesityl is impregnated with elerine, a sort of sub-radical is generated, C<sup>6</sup> H<sup>2</sup>, having a relation to mean, C<sup>6</sup> H<sup>2</sup>, analogous to that which acetyl (3093), C<sup>6</sup> H<sup>2</sup>, has to ethyl, C<sup>6</sup> With the sub-radical thus generated, which was named by its decorate, Kane, ptelevie, the chlorine combines, forming a chloride.

of acctone, and one part of furning nature acid, are mingled, a violent rate tion ensues. After the resulting aggregate has become cool, the addams of water causes the separation of a mixture resembling a vellow oil, which consists of two liquids. The more fluid of these has received the mans of nitrite of the exide of peteleyle. It is heavier than water, and is decomposed thereby. Paper imbued with it burns like prepared tinder. It is capable of bearing the heat of 212° without decomposition, but at a higher temperature it explodes violently; hence it cannot be distilled.

5723. Mesitic Aldehyde, C. HO + HO, or the Hydrated Oxide of

Prolegie.—Of the oily mixture, above described as resulting from the reaction of fuming nitric acid with acetone, mesitic aldehyde forms the more incid portion. It may also be produced by subjecting a mixture of mesityl and mitric acid to ebullition as long as any reaction takes place. From the remula of this, above given, it is seen that it resembles aldehyde in being hydrated oxide of a sub-radical, obtained from another radical by dehyrogenation.

5724. Mesitic aldehyde is a heavy, viscid, reddish-yellow liquid, with a

wectish taste and penetrating odour.

### Of Amyl Ethers.

othetical compound radical, amyl (4023), that, in certain ompounds it has been inferred to play a part analogous that which ethyl, methyl, and other bodies of like kind lay in certain other compounds; and that, especially, it is been inferred that the oxide of this radical exists in he oil of potato spirit, in union with water. This oil beng, therefore, a hydrated oxide of amyl, plays in the computates of amyl, a part like that which alcohol performs in thyl series, or wood spirit in the methyl series. Yet as the oxide of amyl has not been isolated, we have no amylic congener of ether, of which the preparation and properties, in an isolated state, are to be described. Hence, the first object to be presented to the attention of the student is the hydrated oxide.

# Of the Hydrated Oxide of Amyl, or Oil of Potato Spirit, or Amylic Alcohol, C<sup>10</sup> H<sup>11</sup>O + HO.

name has been given, is generated during the vinous fermentation of an infusion of potatoes; and comes over towards the close of the distillation, by which potato spirit is separated from the rest of the products or residue of that process, rendering the water, which simultaneously condenses, milky. Being insoluble in this liquid, it subsides after some time, together with a portion of moisture and alcohol. From the latter of these impurities, it is separated by agitation with water, and from water by chloride of calcium and redistillation. To bring over the pure hydrated oxide of amyl, a temperature of 270° is required.

5727. Potato spirit is a colourless liquid, oily in appearance, with a strong smell, which at first is pleasant, but becomes afterwards extremely nauseous. The inhalation

of the vapour causes asthmatic pains, cough, and em in miting. Its taste is very acrid. It burns with a blood white flame; boils at 270°; has a specific gravity of out at 60°. The density of the vapour is = 3.147, represening four volumes. At 4° it solidifies, forming crystalian plates. It produces a stain on paper, which disposate after a short time; dissolves sparingly in water, to the it communicates its odour; and is miscible in all proper tions with alcohol, ether, fixed and volatile oils, and small acetic acid; dissolves sulphur, phosphorus and iodne. out being altered by them; and may also be mired with solution of caustic potash, or soda, without change: when heated with dry potash, hydrogen is disengaged and valerianate of potash is formed (Dumas, Stas). It about a large quantity of chlorohydric acid gas, with evolution of heat. When mixed with sulphuric acid, a violet coles appears, and the bisulphate of oxide of amyl is product When distilled with dry phosphoric acid, a carbobying is obtained, to which Cahours has given the name des Amylic alcohol combines with the bichloride of forming a crystalline compound, which in the air, and more rapidly when in contact with water, is slowly !solved into its component parts, bichloride of tin and by drated oxide of amyl. See valerianic acid (5283)

5728. Acetate of Oxide of Amyl, Amylo Acetic Ether, Collection Halo Os = Ayl O, Ac Os.—It is easily obtained by distilling a matter of two parts of acetate of potash, one of hydrated oxide of amyleto are of oil of vitriol. The product, after being dried by means of chlorde of all cium, and rectified along with oxide of lend, yields the acetate of purity. It is a colourless liquid, having an ethereal, aromate older, and insoluble in water. It boils at 245°.

Of the Bromide and Iodide of Amyl.

alcohol, and one part of phosphorus, a bromide of amyl, having properties, has been obtained; and likewise an rodide, by the same parts substituting iodine for bromine. The rodide is described as a liquid.

5730. Of Glyceryl and Cetyl, there are no etheres pounds. Accordingly, I here terminate the chapter at ethers.

<sup>\*</sup> A congener of sulphovinic acid (5297).

### OF ANIMAL SUBSTANCES.

5731. Respecting the substances which come under the eceding definition, I had prepared selections from the rmer edition of my Compendium, and from all the other ore recent sources of information within my reach, when e concluding part of the work on Organic Chemistry, by iebig and Gregory, fell into my hands. Finding it to be afficiently condensed, I have concluded to substitute a prion of that work for the matter which I had prepared. Ly only motive for publishing a text book, has been my ability to find any work comprising descriptions of my paratus and peculiar experimental illustrations, and a ving the requisite arrangement and condensation. But the portion of the work by Liebig and Gregory, to hich I have alluded, is deficient neither in arrangement or in brevity, I deem it judicious to embody it in this ompendium.

5732. As certain facts and hypotheses adduced or sanconed by the philosopher of Giessen and his disciples, contitute the organic chemistry now in vogue, and have an nportant bearing on physiology, it seems expedient, so are as practicable, to allow them to be studied in that auhentic form which they have been made to assume by regory, the associate editor of Liebig. Unfortunately, he organic chemistry of Liebig, as translated by Gregory, in general too voluminous and abstruse, to serve as a

hemical text book for a medical class.

5733. I shall change some names in order to produce in accordance with the nomenclature adopted in this work, and to correct an inconsistency in using the noun introgen," and yet employing the adjective "azotized." Evidently either azot and azotized, or nitrogen and nitrogenized, are required by consistency.

The reader will not be misled by some slight differences in orthography, as in Sbrin and fibrine, legumen and legumine.

## Indifferent Nitrogeniaed Substances common to the Vegetable and Anal Kingdoms-Proteine and its Modifications.

5734. "Under this head we have to consider a few very important appounds, which are formed in the vegetable kingdom, and are also must be constitute a large proportion of the animal body. These are Albara, Fibrine, and Caseine. Till very recently, it was believed that vegetable burnen and fibrine differed from animal albumen, fibrine, and cases we the recent researches of Mulder have shown this opinion to be created and Liebig has demonstrated that caseine exists in vegetables with a characters of that found in milk (5023).

5735. "The most important step recently made in advance in the portant investigation is doubtless the discovery, made by Mulder, that also men, fibrine, and caseine, are all modifications of one compound, to with he has given the name of Proteine (from \*guriou\*, I take the first place. I being the original matter from which all these varieties are derived.

5736. "Proteine is composed of carbon, hydrogen, introgen, and open; and Mulder has shown, that two analyses of proteine do not the more than analyses of fibrine, albumen, or caseine do, either from the ther, or from that of proteine, as far as regards these elements. It is further shown, that all of these bodies, whether they contain proteine, however, contains no inorganic matter, albumen, fibrine, at a seine each contain small but essential quantities of mineral substance, as sulphur, phosphorus, potash, soda, common salt, and phosphore of its Purther, it has been established by the still more recent researches of the ochool of Giessen, that animal and vegetable albumen, animal and term ble fibrine, and animal and vegetable caseine, are respectively identate every particular. We may therefore assume that there is but one almost one fibrine, and one caseine; and it is convenient to consider the almost compounds of proteine with small proportions of morganic matter.

5737. "Proteine.—When animal or vegetable albumen, fibras, or a seine, are dissolved in a moderately strong solution of caustic posses, and the solution heated for some time to 120, the addition of acctic and cause a gelatinous precipitate, which has the same composition and possess from whichever of these compounds it has been prepared. When we

washed and dried, this is proteine.

5738. "It forms a yellowish brittle mass, insoluble in water and state. Mulder has analyzed proteine from animal and vegetable a bumen, for fibrine and from cheese, or caseine; and Scherer has analyzed professor animal albumen and fibrine, from the crystalline lens, from has the from born. The results from all these analyses agree best with the mula C\*\* H\*\* N\*\* O\*\* (Liebig); Mulder first gave the formula C\*\* H\*\* N\*\* O\*\*. The symbol of proteins is Pr.

5739. "Proteine combines with both acids and bases: with diluxed sephuric acid it forms sulphoproteic acid, Pr + SO<sup>3</sup>; with diluted chlorely-dric acid, another acid, Pr + 2HCl. When obtaine is passed through a solution containing proteine, white flocks are deposited, which Mulder call

chloroproteic acid, Pr + ClO<sup>a</sup>. (Mulder.)

5740. "When proteine, or any of its modifications, is digested in mine acid, a yellow compound is formed, along with oxalic acid and ammune. The yellow compound is called xanthoproteic acid, and its formula so

Nº O19, 2HO. It seems to combine both with acids and bases. Its

with bases dissolve with a red colour. (Mulder.)

741. "When boiled with an excess of caustic potash, proteine, albumen, are decomposed, yielding, besides ammonia and carbonic and formic s, three azotized products, protide, erythroprotide, and leucine. Eryprotide is a reddish-brown amorphous mass; and its formula, in the pound it forms with oxide of lead, is said to be C<sup>18</sup> H<sup>8</sup> NO<sup>5</sup>. Protide yellowish, soluble, uncrystallizable substance, and its formula is C<sup>18</sup> H<sup>9</sup>. Leucine crystallizes in shining scales, which sublime unaltered at '. Its formula is C<sup>18</sup> H<sup>19</sup> NO<sup>4</sup>. With one atom of hydrated nitric it yields nitroleucic acid, which forms crystallizable salts. Leucine also be formed by the action of sulphuric acid on proteine or its com-

742. "According to Mulder, proteine combines with the oxides of lead silver in the proportion of ten atoms to one of the base: and the same ant of proteine is contained in albumen and fibrine; the former being

 $+ S^4 + P_2^4$ , the latter  $10Pr + S^4 + P_2^4$ .

"According to Liebig, (Animal Chemistry, p. 106,) proteine is uced by vegetables alone, and cannot be formed by animals; although unimal organism possesses the power of converting one modification of sine into another, fibrine into albumen, or vice versa, or both into cauche, &c. In this point of view the vegetable forms of proteine, vegetable nen, fibrine, and caseine, become signally important, as the only ces of proteine for animal life, and consequently of nutrition strictly so d, that is, the growth in mass of the animal body.

744. "Proteine is never found, as such, in nature; but occurs in the e of albumen, fibrine, or caseine, both in vegetables and animals, and

me other forms in the animal body.

### Modifications of Proteine.

45. "Albumen.—This important substance forms the white of eggs, occurs in large quantity in the blood. It is also found in other animal, and in most of the animal solids.

'46. "It occurs also in many vegetable juices, and in many seeds, such its, almonds, &c. From whatever source it is obtained, its properties

he same.

ids. (Mulder.)

47. "Albumen is naturally soluble in water, and is found dissolved in erum of the blood, and in vegetable juices. The white of eggs is quite le; and the albumen of wheat flour also dissolves in water, if it have purified without the application of heat. But when it has once been d to 160°, it becomes insoluble; and, if previously dissolved, a heat of causes the dissolved albumen to coagulate, and the coagulum is insoin water. Hence albumen is described in two states, the soluble and pagulated.

'48. "If white of egg, or serum of blood, be dried up at 120°, the resisoluble albumen in an impure state. It may be purified by being washed with cold ether and alcohol, which remove fat, salts, and other

gn matters.

49. "Dry soluble albumen, when placed in water, first swells up, and forms a glairy fluid. This solution is coagulated by heat, by acids, leohol, by creosote, &c. The acids which do not coagulate albument ectic acid, phosphoric and pyrophosphoric acids. The coagulated al-

burnen dissolves, with the aid of heat, in strong hydrochloric acid, paducing a purple solution. This reaction takes place with all the motion tions of proteine, and indicates a great similarity of constitution among them.

5750. "The solution of albumen is also congulated or precipitated by the acetate of lead and the bichloride of mercury, and by infusion of galls; also by the ferrocyanide of potassium if acetic acid be added. From the insolubility of the precipitate with bichloride of mercury, white of egg, but up with water, is used as an antidote to that poison. One egg combine with about four grains of corrosive sublimate.

5751. "The precipitates formed by acids are compounds of allumes with the acid employed. They are soluble in pure water, but quite units

ble in diluted acids.

5752. "Congulated albumen is quite insoluble in water, but is ready dissolved by caustic alkalies, which it even neutralizes. These component

yield insoluble albuminates with the metallic salts.

from one to two per cent. of phosphate of lime; and soluble altuments pears to possess the property of dissolving that salt, a property which embles the blood to convey to the bones their earthy part, and protects the

to carry away that which is found in the urine.

5754. "When albumen is analyzed, it yields the same results as proteine, in regard to carbon, hydrogen, nitrogen, and oxygen; but it commutes than one per cent. of sulphur and phosphorus together, which are absent in proteine. According to Mulder, it is 10Pr + S<sup>5</sup> + P½; but we have no means of determining with accuracy such small proportions of subbar and phosphorus, and it is therefore preferable to represent albumen as proteine with certain small indeterminate proportions of sulphur and phosphorus. When burned, it also leaves ashes, which contain phosphorus lime and alkaline salts.

5755. "To prove the presence of unoxidized sulphur in albume, in solve it in potash, then add acetate of lead as long as the precipitate forced is redissolved, and heat the solution to the boiling point. It instants becomes black by the separation of sulphuret (sulphide) of lead. The sortest applies to fibrine and caseine. (Liebig.) It is not known in what the phosphorus exists in albumen, after phosphate of lime has been apprated. The fetid smell of putrefying albumen, indicates distinctly the persence of sulphuretted hydrogen (sulphydric acid) among the products of a putrefaction.

5756. "When the juice of many vegetables, after being separated from the coagulum or deposit which spontaneously forms in it, and what a vegetable fibrine, is heated, a new coagulum is formed, which is regestable albumen. When nuts, almonds, and similar seeds, are freed from the clabs oil by pressure, the residue is chiefly vegetable albumen in the whole form. It is, in every respect, identical with the albumen of eggs and a

blood.

5757. "Albumen must be considered as the true starting point of all the animal tissues. This appears from the phenomena of incubation, where a the tissues are derived from the albumen of the white and of the volk, which contains albumen also, with the aid only of the air, of the oily matter of the yolk, and of a certain proportion of iron, also found in the yolk. It is dear from this, that albumen may pass into fibrine, caseine, membranes, horehair, feathers, &c.

5758. "Fibrine.—This modification of proteine occurs, like albumen, in two forms, dissolved and coagulated. The former is found in fresh-drawn blood and in fresh-drawn vegetable juices, from both of which it coagulates spontaneously on standing. In the coagulated state it is found in muscular fibre, and in the gluten of wheat flour and the seeds of the cerealia generally.

5759. "The characters of insoluble or coagulated fibrine closely resemble those of coagulated albumen. With strong acetic acid it forms a jelly, which may be dissolved by boiling water, and a precipitated by ferrocyanide of potassium. It is similarly acted on by other acids; and, like albumen, dissolves in alkalies, which it neutralizes. It gives a purple solution

with strong hydrochloric acid.

5780. "When fresh blood is allowed to stand, the fibrine dissolved in it coagulates very soon, and forms the clot, which, however, is coloured by the globules of the blood; but if the blood be stirred with a stick while coagulating, the fibrine adheres to the stick in grey stringy masses, which dry, like albumen, into a horny matter. Like albumen, it contains sulphur and phosphorus, and its ashes contain phosphate of lime. It contains less

sulphur, however, then albumen.

5761. "As albumen, during incubation, passes into fibrine, so fibrine, in the animal body, passes into albumen; for example, in the case of an animal fed on muscular fibre, whose blood contains the usual proportion of albumen. Nay, Denis has shown that the fibrine of venous blood, by digestion with a solution of nitre, is dissolved, and acquires the characters of albumen, being coagulated by heat and by acids. Scherer has shown that the fibrine of arterial blood does not undergo this change, nor that of the buffy coat, nor even venous fibrine after exposure to the air for some time. Hence he concludes that it is rendered incapable of dissolving by the action of oxygen, and that the fibrine of venous and of arterial blood are thus distinct; the former being soluble, the latter congulated.

5762. "He found that the above mentioned solution of venous fibrine in nitre, when exposed to the air, deposited an insoluble matter, identical with arterial fibrine. He also observed, that, after being boiled for a short time, venous fibrine became insoluble, and had lost the property, possessed by it when fresh, of absorbing oxygen and giving off carbonic acid.

5763. "The fibrine which spontaneously coagulates from certain vegetable juices, such as those of carrots, turnips, and best-root, and that contained in the gluten of wheat flour, are identical in properties and composition with animal fibrine.

5764. "Fibrine, both animal and vegetable, is a most important element of nutrition, and yields, in the animal body, albumen, caseine, and the tissues derived from them.

5765. "Caseine.—This, the third important modification of proteine, is found in milk, and constitutes that ingredient which is neither coagulated spontaneously, like fibrine; nor by heat, like albumen; but by the action

<sup>\*</sup> To determine the quantity of fibrine in blood, M. Simon receives it in a flask containing little bits of metal first weighed without the flask, then with this recipient; and after the introduction of the blood, the flask and its contents are weighed together. On agitating the flask, the bits of metal become conted with fibrine. Subsequently they are washed with water, dried, and weighed. By these means, guarding against any deficit, the weight of the fibrine may be deduced. Agreeably to the same authority, menstrual blood contains no fibrine "Berselius' Report, 1841, page 263.

of acids alone. Cheese, made from akimmed milk and well preset, a nearly pure caseine. A substance quite identical is found abundantly at the seeds of leguminous plants, and was formerly called Legumino. Ling has recently shown that legumine is nothing but caseine; and, from whether source, it is found to be a compound of proteine. Thus its assign gives the same results as those of fibrine and albumen for the four organs elements; and it differs from these bodies in containing no free phopiaris. Its ashes are very rich in phosphate of lime and in potash; and a the point also animal and vegetable caseine agree.

5766. "Coagulated caseine is generally a compound of caseine was the acid employed to coagulate it. When milk, on standing long, coagulate it is found to contain free lactic acid, some of which by combining with the caseine has caused the precipitate. When sulphuric acid is used, the caseine is sulphate of caseine; which, when the acid is removed by curious

ate of lead, yields pure caseins.

dissolved by water, and never forms a clear solution. It is not ready acetic acid, but in other respects resembles a solution of albumen, except the it is not coagulated by heat. When milk is placed in contact with read, which is the lining membrane of a calf's stomach, it is coagulated. Laby has shown, that, unless the membrane be in a state of decomposition, the change does not take place; and it probably depends on the formation and the fermenting influence of the membrane, of sufficient lactic acid water tralise the alkali of the milk, and thus coagulate the caseine. When seek milk or cream is used, the cheese contains much butter besides the casein.

5768. "When milk is heated in an open vessel, a pellicle is formed, which if removed, is continually renewed, and is insoluble. It is owing to to action of oxygen, for it does not form in an atmosphere of carbonic and

(Scherer.)

5769. When peas, beans, or lentile are softened in cold water, the ground with that fluid, and the mass further diluted, and strained through a fine sieve, there passes through a solution of caseine in which such suspended. When the starch has settled, the supernatant liquid a sattern of caseine, which is always, like milk, turbid, partly from suspends fat, partly from the gradual action of the air on the dissolved caseine, lack acid being slowly formed, which causes a gradual separation.

5770. "This solution has all the characters of skimmed milk; it is congrelated by acids, not by heat, and forms a pellicle when heated. It also engulates after long standing from the formation of lactic acid; and, when the congulum putrefies, the odour is exactly that of putrid cheese. Least.

5771. "The ashes of soluble caseine, whether animal or vegetable, we very strongly alkaline; and there is reason to beneve that the point found in the ashes had served, by combining with the caseine, to reader a soluble.

5772. "Caseine occurs also in the only seeds, such as almonds, nuts, be along with albumen, and must be considered as a very important element of nutrition.

5773. "Scherer, by acting on the serum of blood with water and a limit caustic potash, obtained a neutral solution, which no longer congulated by heat, but formed a pellicle like milk. As this pellicle appeared identification with that from milk, the experiment seems to prove the conversion of allowers into caseine.

5774. "Mulder considers caseine to be 10Pr + S; but pure caseine s

own, and caseine, as it usually occurs, contains 6.5 per cent. of incomatter, chiefly phosphate of lime and potash. There is no doubt, er, that the organic elements of caseine are united in the same pro
as those of proteine, albumen, and fibrine; while, like the two latter, is a purple solution when heated with strong chlorohydric acid. The of milk, also, in the nutrition of young animals proves that caseine the of conversion into albumen and fibrine; while the production of an animal fed on albumen or fibrine, or both, shows that these boasy be reconverted into caseine.

5. "We may exhibit the connection between these substances as Pr represents proteine, C\*\*\*H\*\*\*N\*\*O\*\*\*. P and S represent, not lents, but only small indeterminate quantities, of phosphorus and

r.

Albumen is . Pr+S\*+P+salts. Fibrine is . Pr+S +P+salts. Caseine is . Pr+S +salts.

8. "We can thus easily understand the formation of any one of them proteine, or the conversion of one into the other. Albumen, losing sulphur, becomes fibrine; and fibrine, losing its phosphorus, becaseine: but the salts are not exactly the same, nor in the same proin all the three cases.

#### The Blood.

7. "This important fluid, from which all parts of the body are formsenses very remarkable properties. In the veins it is dark-coloured,
arteries bright red. When drawn, it presently forms a red clot,
sed, as we have seen, partly of fibrine, while the serum contains a

uantity of albumen.

\* The colour of the clot is owing to a compound which has been \*Hematorine\*, which has many properties in common with albumen; globules of the blood, in which the colour naturally resides, are not ed of hæmatosine alone, but contain another albuminous compound, the name of \*Globuline\* has been given. It is probable that neither

s compounds is known in a state of purity.

t. "To obtain them, blood is well stirred to separate the fibrine, and with six volumes of a saturated solution of sulphate of soda, in which bules are insoluble. They are then boiled with alcohol acidulated loburic acid, which dissolves a sulphate of harmatosine, and leaves a of globuline. The red alcoholic solution is mixed with carbonate nonia, which separates the sulphuric acid as sulphate of ammonia, with a little globuline. The filtered solution, being evaporated, leaves usine as a dark brownish red mass.

"Hæmatosine thus prepared is insoluble in water, alcohol, and out forms red solutions with alcohol, to which either acids or alkalies ied. Its ashes contain iron, but Liebig and Scherer have shown that colour does not depend on that metal, which may be removed either e globules, or from hæmatosine, by strong sulphuric acid, without ing the red colour; and in this experiment the red matter left gives a sh, free from iron. Iron, however, is essential to the blood, and is sently supplied in the food. The ashes of almost all vegetables contitle iron; flesh, of course, does so, as it is mixed with blood; and t of egg is found to contain an oily matter, of which iron is an interest.

5781. "Globuline forms the principal part of the blood-globules. It has not been obtained in a pure state, but has all the characters, as well in the composition, of dissolved albumen. The compound with sulphunc and above mentioned, is grey, or white, and was found to contain four man

of sulphuric acid and one of proteine.

matters. When heated, the albumen coagulates, and floats in a warry liquid called the Serosity. This contains common salt, sulphan prophetes, and carbonates. The blood probably also contains the ingulates of the secretions and excretions, such as bile and urine; but there are a small a proportion, that, except in cases of disease, it is hardly possess detect them. The fatty matter in blood is obtained by drying up the sease and digesting with other. It consists of the usual animal fats, and a seal likewise to contain cholesterine, or the fat of bile.

5783. "The following table gives the results of two careful analysis of

blood by Lecanu:

|                   |                      |                     |         |                       |               |       |       | Home     | n blook  |
|-------------------|----------------------|---------------------|---------|-----------------------|---------------|-------|-------|----------|----------|
| Water             |                      | -                   |         |                       |               | -     | -     | 780 145  | 795.500  |
| Fibrine           | -                    | -                   | -       |                       | -             | -     | -     | 2 100    | 3.565    |
| Colouring         | matter               | (hemate             | bette 4 | nd glob               | uline)        |       | -     | 133 000  | 117,60   |
| Albumen           | -                    |                     |         | -                     |               |       | -     | 65-090   | (8.41)   |
| Crystallin        | e fat                |                     |         | -                     | -             |       | -     | 2.430    | 4300     |
| Oily matte        |                      | -                   | -       | -                     |               | -     | -     | 1 310    | 250      |
| Extractive        | malter               | (soluble            | in wa   | ter and               | alcohol)      | -     | -     | 1 790    | 1,500    |
| Albuminat         |                      |                     | -       |                       | -             | -     | -     | 1.985    | 2,000    |
| Alkaline c        |                      |                     |         |                       |               |       |       | 8.370    | 126      |
| Carbonate<br>magn | s of hir<br>caia, an | ne and<br>d iron, j | megno   | min, pho<br>to of iro | osphates<br>n | of li | me, } | 2100     | 1.64     |
| Loss              | -                    | -                   | -       |                       | -             | -     | -     | 2.400    | 2.386    |
|                   |                      |                     |         |                       |               |       |       |          |          |
|                   |                      |                     |         |                       |               |       |       | 1000.000 | 1000.000 |
|                   |                      |                     |         |                       |               |       |       |          |          |

of proteine, its composition cannot be very different from that of process of its modifications. In fact, dried blood, when analyzed, yields the number C<sup>48</sup> H<sup>39</sup> N<sup>6</sup> O<sup>15</sup>, which is proteine, C<sup>48</sup> H<sup>36</sup> N<sup>6</sup> O<sup>15</sup> + HO + H<sup>3</sup>. (Paying and Boeckmann.) This excess of hydrogen is probably derived from the presence of fat.

5785. "From the blood, that is, from the compounds of proteon as blood, are derived all the animal tissues. Some of these are compounds; but a proteine, others have no longer the characters of such compounds; but a

all cases they are derived from proteine.

5786. "Muscular tissue, or muscular fibre, is composed chedy of fibrine, mixed however in the ordinary state with blood, membrane, we your matter, and fat. Dried flesh, when analyzed, gave the same forms as dried blood, namely, (\*\*\* H\*\*\* N\*\*\* (Playtair and Boeckmann.)

5787. "When flesh is acted on by hot water, there is dissolved a paintity equal to 17 per cent, of its weight. The dissolved matter contains the salts of flesh, and several organic matters probably produced in the operation, the nature of which is very little known. One of these has been described under the name of Osmazome, and is supposed to give to soup and dressed ment their peculiar flavour: but osmazome is certainly not a pure substance, as at present known; and the whole subject of the changes

<sup>&</sup>quot;In other words, comesome to not a definite compound to which any formula or be assigned, but probably a mixture of substances not yet distinguished.

moduced in food by cooking is understood to be under investigation by liebig.

5788. "Gelatinous tissue.—Under this name are included the organic same of the bones, that of tendons and ligaments, the cellular tissue, the kim, and the serous membranes. All these substances dissolve by long-cantinued boiling in water, and the solution on cooling forms a jelly. The same forms of gelatine, from hoofs, hides, &c. are called Glue; that from hand finer membranes is known as Size; and the purest gelatine, from air-bladders and other membranes of fish, is called Isinglass. Gelatine not exist as such in the animal tissues, but is formed by the action of pailing water."

5769. "Gelatine is soluble in water, and the hot concentrated solution water a jelly on cooling. It is precipitated by tannic acid, forming an insomble compound, which forms the chief part of leather. Leather is made y steeping softened skins in a strong infusion of oak-bark, catechu, or their astringent vegetables containing tannic acid. Skins are prepared in their ways, yielding different kinds of leather, such as tawed leather, washanther, &c.; but the process of tanning depends on the action of tannic

cid on the gelatinous tissue.

5790. "Gelatine, when acted on by sulphuric acid, yields gelatine suuse, or glycicoll, C° H<sup>7</sup> N° O<sup>5</sup>, 2HO. When treated with potash, it is
nice to yield glycicoll and leucine. Glycicoll unites with oxide of lead,
arming a compound, C° H<sup>7</sup> N° O<sup>5</sup>, 2PbO. (Mulder.) It also combines with
itric acid, forming a compound acid, C° H<sup>7</sup> N° O<sup>5</sup> + 2NO<sup>5</sup> + 4HO, which
rystallizes, and forms double salts with bases. That with lime is CaO,C°
17N°O<sup>5</sup> + 2(CaO, NO<sup>5</sup>).

5791. "According to Scherer, the composition of gelatinous tissue is expresented by the formula C<sup>48</sup> H<sup>41</sup> N<sup>7</sup> A O<sup>18</sup>, or doubled, C<sup>66</sup> H<sup>68</sup> N<sup>15</sup> O<sup>26</sup>; which latter formula represents 2 atoms proteine + 3NH<sup>8</sup>, +HO + O<sup>7</sup>.

5792. "Although gelatine is thus nearly related to proteine, and is doubtses formed from one or other of its modifications, yet it has none of the
haracters of a compound of proteine. It does not yield proteine when
cted on by potash, and it does not produce a purple colour with hydrochlo-

ic acid. It therefore no longer contains proteine.

5793. "This accounts for the fact, that animals, fed exclusively on gelaine, die with the symptoms of starvation. The gelatine, containing no proteine, cannot yield albumen, fibrine, or caseine; and it has already been tated that the animal system, although it can convert one form of proteine another, cannot form proteine from compounds which do not contain Blood therefore cannot be made from gelatine, and the animal soon lies. But when mixed with other food, especially compounds of proteine, platine may be useful, and may serve directly to nourish the gelatinous insues. (Liebig, Animal Chemistry, 98, 130.) This would explain the set of gelatine as a part of the food of convalescents, whose debilitated system cannot readily convert albumen, &c. into gelatine for the nutrition of hese tissues, and finds it ready-made in the food. The experiments of D'Arcet on the gelatine from bones have proved, that, as part of the diet in sospitals, gelatine produces the best effects, and materially abridges the

How happens it then, that, as it exists in the hides or skins of animals, it combines with tannic acid to form a substance (leather), precisely the same in its chemical composition as the precipitate formed by a solution of gelstine with that acid?

period of convalescence. When it is given alone, all animals soon t

it with disgust, and die if confined to gelatinous food.

5794. "Chondrine.—This substance forms the tissue of cartilego occurs in the ribs, traches, nose, &c. and of the comes. It is slowly solved by boiling with water, and when dry resembles glue. But it from gelatine in not being precipitated by tannic acid, and in giving cipitates with acetic acid, alum, green vitriol, and acetate of lead. If when in the cartilaginous state, are composed of it. According to but it is composed of C40 H40 N6 O20; that is, of proteins + 4H0 + Chondrine leaves, when burned, from 4 to 6 per cent. of substant bone-earth.

5795. "Arterial Membrane.—The middle coat of the artery, what very elastic membrane, leaves, when burned, 1.7 per cent of According to Scherer, it is composed of C\*\* H\*\* N\*\* O\*\*\*; that is, per + 2HO.

5796. "Horny matter.—This occurs in two forms, membrane compact. The former constitutes the epidermis, and the epithelium, lining membrane of the vessels, of the intestines, and of the pulmonary. The latter forms hair, horn, nails, &c.

5797. "Scherer has analyzed numerous specimens of both his horny matter, and deduces from his results the formula C\*\* H\*\* N

that is, proteins  $+ NH^s + O^s$ .

5798. "Horny matter, when acted on by potash, yields proteins

addition of acetic acid.

5799. "Feathers are closely allied to horny matters, but, according to the second scherer, contain one atom of oxygen less; the formula of feathers, in from his analysis, being C<sup>46</sup> H<sup>59</sup> N<sup>7</sup> O<sup>16</sup>.

5800. "Pigmentum nigrum oculi. - This substance, according to St contains more carbon than any of the preceding; but its formula is

been ascertained.

5801. "It is to be particularly observed, that the formulæ above for the principal tissues of the body, are only intended to show the rethey actually bear to proteine. It is not meant that they are formed a body by the addition of water, ammonia, or oxygen, to proteine: o contrary, we are as yet ignorant of the conditions under which the produced; and in some cases, as, for example, in gelatine, several directors may be taken of their formation.

#### Brain and Nervous Matter.

5802. "Nervous matter is distinct from all other animal tissues, a produced by the animal system exclusively. In composition it is mediate between fat and the compounds of proteine, containing natrogen, is absent in fats, but in far smaller quantity than proteine does: and on the other hand, much richer in carbon than proteine or its compound is appears likewise to contain phosphorus as an essential ingredient.

5803. "From the recent researches of Fremy, brain appears to e a peculiar acid, analogous to the fatty acids, which he calls cerebre and which contains a trogen and phosphoras: this is mixed with an minous substance, with an only acid—the oleophosphoric acid, with a terine, and finally with small quantities of oleine and margarine. a oleic and margaric acids.

5804. "The two acids peculiar to the brain and nervous matter.

torretimes free, but generally combined with sods or with phosphate of lime.

\$605. "Cerebric acid is extracted by other from the brain after it has been exposed to the action of boiling alcohol, which congulates the albumenths matter deposited on cooling by the other is a mixture of cerebric acid, penerally combined with soda or bone-earth, eleophosphate of soda, and a little albumen.

2.5806. "This mixture is acted on by alcohol, acidulated with sulphuric said, which precipitates sulphates of lime and soda, and albumen. The filegred solution contains cerebric and eleophosphoric acids; cold ether resolves the latter, and the former is purified by solution in hot ether and anystallization.

5807. "When pure, it is white, crystalline, and pulverizable. In hot water it swells up like starch, but does not dissolve. It contains phosphorus, but no sulphur if purified from albumen; the phosphorus amounts to barely one per cent.; and it contains 2.3 per cent. of nitrogen. It has the characters of a fatty acid, but its acid properties are feebly marked.

with the alkalies it forms soaps, and its compound with soda appears to exist in the brain. When it is long boiled with water or alcohol, it is resolved into oleine and phosphoric acid. This change is accelerated by acids, but it takes place also spontaneously at the ordinary temperature, only more slowly; and the presence of animal matter in a state of decomposition seems to cause it to be resolved into oleine and phosphoric acid. Thus, when brain has been allowed to undergo partial putrefaction, it no onger yields oleophosphoric acid, but oleine and phosphoric acid. It consine two per cent, of phosphorus. The oleine of this acid is identical with that of burean fat.

5809. "Cholesterine.—This fat, as extracted from the brain, in which t occurs in considerable quantity, has the same composition and properties the fat of biliary ca.culi. (Couerbe; Freiny.) Fremy has also succeeded in detecting in the liver, traces of the characteristic fat acids of the train.

5810. "The grey portions of the brain appear to be chiefly albuminous; while the white portions consist of an albuminous tissue similar to the grey, but loaded with the fats above described.

5811. "The softening of the brain in diseases of that organ seems to be he result of putrefaction, and is accompanied by the separation of the cleine from the phosphoric acid. The cleine itself also is decomposed, yielding free cleic acid.

5912. "There can be no doubt that the brain and nervous matter (which squite similar to brain) are formed in the body from compounds of protaine, either by the loss of some azotized compounds, or by the addition of highly carbonized products, such as fat. But we are ignorant in what part of the body, or by what organs, nervous matter is prepared. This point requires minute investigation. In the mean time, according to Chevreul, the fatty matters, which occur in small quantity in the blood, are similar to those of the brain.

5813. "Bones.—The bones of animals are composed of bone-earth and polatinous tissue. By the action of hydrochloric acid, the earthy matter is bone, and the animal tissue is left. It is soft, retains the form of the bone, and, when dried, becomes brittle and semitransparent. When boiled

with water, it yields a solution of gelatize, fatty matter remaining unisolved.

5814. "The earthy matter is formed of a peculiar phosphate of lim, 8CaO + 8P° O° = HO, 2CaO, P° O° + 2(8CaO, P° O°); that is, a mappound of two forms of tribasic phosphate. It forms rather more than hill the weight of the bone, and contains a variable proportion of carbonized lime. Fluoride of calcium is sometimes, but not always, present a recent bones; in fossil bones, and in human bones from Herculaneum, it a shape found.

5615. "Testà contain the same ingredients as bonce, but the properties of earthy matter is greater, amounting to nearly seventy per cent. The enumed of the teeth contains no animal matter, and fluoride of calcum a found in it.

5816. "In rickets, the proportion of earthy matter is much dismaint. Callus and excetosis are said by Valentin to contain more carbonate of law than sound bone, and carious bone to contain less.

5817. "When bones are heated in the open fire, they leave an early skeleton, which is quite white, and has the form of the bone. If bone are heated in close vessels, they give off carbonate of ammonia and tarry products, and leave a black mass, which consists of bone-earth, with about per cent. of finely divided charcoal. It is called bone or ivory-black, and is much used to decolorize organic solutions.

#### Animal Secretions and Excretions.

5818. "Milk.—This important secretion, destined for the support of the young of the mammalia, is characterized by the caseine it contains. But it also contains certain oily or fatty matters which constitute butter, and which, besides fats analogous to the ordinary animal fats, contain certain volatile acids (5052, 5056), to which the smell and peculiar taste of butter are owing. Milk further contains sugar of milk, or lacture (4070; and, when the caseine has been coagulated by an acid, the whey, besides incline and salts, contains an albuminous matter, which is coagulated by but.

5819. "The composition of milk is such, that it is capable of supporting animal life without any other food. Its caseine and albumen serve for the formation of blood, and for the nutrition of the animal tissues, while as a gar and fat support respiration; and it furnishes, besides, all the salts when the body requires.

5820. "The following table exhibits the composition of the milk of someon, of the ass, and of the cow. (Henry and Chevallier.)

|                   |   |   |   | Milk of |        |        |  |
|-------------------|---|---|---|---------|--------|--------|--|
|                   |   |   |   | Woman.  | Ass    | Cow    |  |
| Cheese or caseine | ŀ | - | • | 1 52    | 1 82   | 4.48   |  |
| Butter -          | - |   |   | 3.55    | 0 11   | 3.13   |  |
| Sugar of milk     | • | • | • | 6 50    | 6 08   | 4.77   |  |
| Salts and mucus   | - |   |   | 0.45    | 0.34   | 0.60   |  |
| Water -           | • | - | - | 87 98   | 91.65  | 87.02  |  |
|                   |   |   |   | 100.00  | 100 00 | 100.00 |  |
|                   |   |   |   |         |        |        |  |

5821. "When the food is highly farinaceous, the proportion of butter a increased; but when the food contains much of the compounds of protests there is less butter and more caseine present. The more active exercise a taken, the smaller also is the proportion of butter.

5822. "Milk, after it has become sour, undergoes the vincus fermenta-

ine (5215).

5623. "Saliva.—This fluid, secreted by the salivary glands, is comsecond of water, with about one per cent. of solid matter, partly saline. It when contains a trace of sulphocyanide of potassium, or at least of a sait which strikes a red colour with persalts of iron; but this might be done by un acetate. The animal matter of saliva has been described under the same of Balivary matter. It is soluble in water, and not congulated by hant.

- 5824. "Saliva possesses, in an eminent degree, the property of frothing with air, like a solution of soap; and Liebig (Animal Chemistry, p. 118) sonceives that its use is to introduce in this manner, during mastication, a sertain quantity of air into the stomach, the oxygen of which is employed an digestion.
- As I find that Berzelius, Graham, and Kane, give mere importance to peptine han is accorded by Liebig and Gregory, I deem it expedient to subjoin the following abridged account of it, prepared by me recently, from Berzelius' Report, 1840, 193, and Graham's Elements, 1930.

#### Pepsins.

The same of pepsine has been given to a peculiar matter constituting the active stinciple of the gastric fluid, the discovery of which is due to Mr. Wasmann. Pepsine may be obtained by infusing the mucous membrane of the stomach in aciduated water. The solution thus procured, has the property of dissolving the coagu-

sated white of egg completely in half an bour.

When the membrane, without being out into pieces, but well washed, is digested m a large quantity of water, at a temperature between 80° and 90°, a variety of substances are extracted as well as pepsine; but if afterwards cold water be substituted for the warm, scarcely any matter besides pepsine is taken up. The extraction may induce with successive portions of water, until symptoms of putrefaction ensue. The solution thus obtained, with the addition of a little chlorohydric acid, has the property of dissolving coagulated albumen speedily. Pepsine, extracted by these peaces contains a little albumen, which may be precipitated by forecasside of nomeans, contains a little albumen, which may be precipitated by ferrocyanide of po-assrum, or by heating the solution, if not too dilute, to a temperature between 1700 and 2120 without ebuilition. By these means the congulated albumen is precipitated a flocks, with a little modified caseine.

Persine may be precipitated from its solutions by the protosulphate of iron, sul-shate of copper, acetate of lead, or protochloride of tin. From the precipitates thus unde, it may be separated by exposure, while suspended in water, to sulphydric

seid.

In precipitating, pepsine retains a sufficient portion of the sold of the saline pre-sipitant to have a decided reaction with litmus, and is highly endowed with its ap-

propriate solvent powers.

Acetate of pepsine may be procured by decomposing, by sulphydric acid, the prespitate made as above suggested, by acetate of lead, evaporating the residual solution to the consistence of syrup, and subjecting it to alcohol. The acetate separates in white flocks, which, by desiccation, acquire the appearance of a gum, and are eadily soluble in water. Of papeine in this form, one part in 60,000 parts of water, with a minute addition of chlorohydric acid, dissolves indurated albumen within

shout six or eight hours.

A similar efficacy is ascribed to the chlorohydrate of pepsine, which may be obtained by precipitating the solution by bichloride of mercury, and subjecting the pre-

similate to the process above described in case of the acetate.

Mr. Wasmann has remarked, that the pepsine obtained from the pig is devoid of the power to coagulate milk, although that of the calf is highly endowed with this

Agreeably to some comparative trials of the solvent powers of dilute chlorohydric seid, without pepsine, and one other portion of the same acid containing this princi-ple, it appeared that the one was endowed with all the solvent powers of the gastrie baid in a high degree, at ordinary temperatures, while the other, under like circum-stances, displayed them only to an insignificant extent; but when the said, without pepsine, was aided by boiling heat, its solvent powers were equal to that of the solu-

5825. "Gastric Juice.—This remarkable fluid seems to contain in any principle capable of accounting for its solvent power. In the capable stomach it is neutral, but during digestion it becomes acid, from the ration of free muratic acid. According to Wasmann and other change it contains a peculiar principle, Pepsine, which has the property of contiing food, and which is obtained by the action of water on the well-water lining membrane of the etomach of the pig. According to Liebig, lowers, pepsine, as a distinct compound, does not exist. The solution of the mag membrane, alightly acidulated with chlorohydric acid, certainly content albumen and fibrine, if kept in contact with them out of the body at the ordinary temperature. But none of these effects take place, union to membrane has been previously exposed to the air, and is in a state of the composition. Hence Liebig ascribes (Animal Chemistry, 100 sept to solvent power of the gastric juice to the gradual decomposition of a mater dissolved from the membrane, aided by the oxygen introduced in the salva-Albumen, dec. when thus in contact with decomposing or ferment ag milter, are rendered soluble by a new arrangement of their particles. The accumulation of free chlorohydric acid, derived, no doubt, from comme salt, at last puts a stop to further change. The whole food is now broads into the form of chyme, an opaque homogeneous fluid, which afterward passes, first into chyle, and finally into perfect blood. In the chyle, is formation of fibrine has already taken place; for, when drawn, a corplates spontaneously, like blood.

5826. "Pancreatic Juice.—The fluid secreted by the pancreas spated into the duodenum, and mixes with the chyme as the latter leaves is stomach. It contains albumen, and, according to some, caseine, and a acid. Its nature, however, is little understood, and its uses at present of

unknown.

### Bile and Biliary Calculi.

5827. "The bile is a yellowish green viscid liquid, secreted by the liver. It has a faint disagreeable smell; and its taste is at first sweet, afterwards bitter and nauseous. Ox bile has been chiefly examined, but that of san and other animals is very similar. The researches of Tiedemann and Gmelin, of Berzelius and Demarçay, have shown that bile may be made to yield a vast number of different compounds, most of which are products of decomposition.

5928. "The bile, according to Demarcay, contains soda in combination with a peculiar acid, choleic acid. When bile is boiled with an excess of chlorohydric acid, it yields ammonia, taurine, and choloidic acid; and when boiled with caustic potash, it yields carbonic acid, ammonia, and

**ch**olic acid.

5829. "Choleic Acid.—When bile is acted on by alcohol, certain impurities are left undissolved. The purified bile gives with acctate of kad a

"Chyle resembles blood in resolving itself into a coagulum, and a liquid like erum, which, according to Dr. Prout, consists partly of albumen, but principally a incipient albumen. The coagulum, according to Vanquelin, is imperfect fibria. Its Brande considers it as more allied to caseous matter.

The opinions of Front and Vauquelin derive support from the consideration that as chyle is destined to become blood, it may be reasonably expected to contain the principal constituents of that liquid, in a state advancing towards maturity. These inferences respecting chyle, made in the former edition of this Compendium, approx to be sanctioned by those of Liebig expressed in the text.

precipitate of choleate of lead, which, when acted on by sulphuretted hy-

drogen, yields choleic acid.

5030. "It forms a yellow spongy mass, soluble in water and alcohol, which has an acid reaction and a bitter taste, and is decomposed by heat. It combines with soda, forming a compound which Demarçay considers as a soap, the solution of which in water has the physical characters of bile. But although this be the case, and although the composition of the cholcic acid appears to be the same as that of the organic part of bile, yet we cantest consider the bile as choleate of soda; for the latter is decomposed by acetic acid, which has no action on bile.

5881. "According to the analyses of Demarcay and Dumas, as calculated by Liebig, the formula of choleic acid is C76 Hee No Oas, and this for-

mula may represent also the organic part of the bile.

5832. "When choleic acid is boiled with chlorohydric acid, it yields ammonia, taurine, and choleidic acid. The latter, being insoluble, is deposited, and the taurine is extracted from the mother liquor by concentrating and adding a large quantity of alcohol, when the taurine slowly crystallizes.

5833. "Choloidia Acid.—This acid is solid, fusible, of a yellow colour and bitter taste, insoluble in water, soluble in alcohol. It combines with bases, neutralizing them, and forming salts which are soluble in alcohol.

It contains no nitrogen, and its formula is C7s Has Oss.

There remains 1 atom choloidic acid

5834. "Taurine.—This substance forms white crystalline needles, which are soluble in water, and sparingly soluble in alcohol. Its formula is C\* H' NO.

5835. "The production of these substances is easily explained.

We subtract 1 atom taurine C4 H7 NON And 1 atom ammonia . C4 H7 NON H2 N C6 H10 Na O10,

5636. "Cholic acid.—This acid is formed, along with carbonic acid and ammonia, when bile or choleic acid is boiled with an excess of caustic potash. It is precipitated by acetic acid, and purified by alcohol from unaltered choleic acid.

5837. "When pure, it forms fine needles, which are permanent in the air; or large tetraedrons, which become opaque on exposure. It is insoluble in water, soluble in alcohol and ether. It forms neutral salts with bases. Its formula is C<sup>74</sup> H<sup>80</sup> O<sup>18</sup>, and its formation is easily explained.

5838. "Berzelius states that the bile is far from being so simple in its constitution as Demarçay supposes; and by a series of ingenious processes has obtained from the bile a number of different substances, which he has named Biline, Biliverdine, Dyslysine, Fellinic Acid, and Cholinic Acid, besides taurine and choic acid, as already described. Biline is essentially the same as Demarçay's choleic acid; and it is probable that most of the others are products of decomposition. But even supposing choleic acid to be composed of two or more different compounds, not isolated by Demar-

pot overlook the fact, that it is constant in its composition, and that fact this composition we can deduce the principal products of the action of the composition we can deduce the principal products of the action of the compound and alkalies on bile. It is choicic acid or bile as a whole, whether a becompound or a mixture of several, to which we have to look for the companion of the changes by which bile may be formed or decomposite. The researches of Berzelius have rendered it probable that choleic acid is single compound; but this does not affect its ultimate composition, for a relation to decomposing agents. It is also clear that bile is very process change in almost all circumstances, and yields a great variety of protest, most of which have little physiological interest. For these reasons we had merely refer the reader to the elaborate paper of Berzelius in the bound in a calculated and interpreted by Liebig, they admit of direct application is physiology.

of soda is dissolved, and the residue is found to contain mucus, and perfectly matter. The latter consist of cholesterine and ordinary fat, and periods contain a portion of the peculiar fats of brain. The dissolved periods besides true bile, contains a small portion of soaps of marganic and dis

acids with soda.

5840. "The sugar of bile or picromel of Gmelin, so called from a season and bitter taste, appears to be choleic acid or biline, altered by the process."

to which it has been subjected.

der, and are often the cause of much suffering, are almost always compant of cholesterine, with more or less colouring matter. Hot alcohol distribution cholesterine, and deposits it in shiring scales on cooling. These calculates

have often a form nearly cubical, and a pearly lustre-

Goebel in a biliary concretion, and appears to be the chief constraint of the concretions called bezoar stones, which occur in herbivorous analyst according to Ettling and Will, its formula is C<sup>40</sup> H<sup>56</sup> O<sup>6</sup>. It is soluble in hot alcohol, and forms a crystalline powder on cooling. It is insoluble in water, and forms with alkalies soluble soaps, with oxides of lead and alver insoluble compounds. It is decomposed by heat, and when acted on by nitric acid yields a new acid.

5943. "According to Liebig, who deduced the above formula from the analysis of Ettling and Will, lithofellic acid may be formed, along with

hippuric acid, by the exidation of choleic acid.

1 eq. choleic acid, C76 Het Ns Osz and 10 eq. oxygen, O10 
$$= \begin{cases} 2 \text{ eq. hippuric acid, C56 Hit Ns Oss} \\ 1 \text{ eq. lithofellic acid, C56 Hit O4} \\ 14 \text{ eq. water,} \end{cases}$$

$$= \begin{cases} 2 \text{ eq. hippuric acid, C56 Hit Ns Oss} \\ 14 \text{ eq. water,} \end{cases}$$

$$= \begin{cases} 2 \text{ eq. hippuric acid, C56 Hit Ns Oss} \\ 14 \text{ eq. water,} \end{cases}$$

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$$= \begin{cases} 2 \text{ eq. hippuric acid, C56 Hit Ns Oss} \\ 14 \text{ eq. water,} \end{cases}$$

5844. "Excrements.—The excrements of man contain about one-fourth of their weight of solid matter. The ashes of dry fæces amount to 13.56 to 15.00 per cent., and are composed of phosphates and other salts. The excrements also contain nitrogen, and yield ammonia when they purefy. The value of night-soil as manure depends on the salts and ammonia of the fæces, and also in a great measure on the ammoniacal and other salts of

rine. The colour of faces is generally said to be owing to bile; but g states that there is only a mere trace of bile, if any, to be found in eces either of man or animals. The yellow matter of faces is insolulated alcohol, with the exception of a small proportion, and even that has be characters of bile.

- 45. "Lymph.—The lymph of cellular membrane is water, with a trace of albumen and of common salt. The lymph secreted by the s membranes is much more highly charged, containing seven or eight ent. of albumen and salts. It coagulates when heated, or by the act nitric acid. The liquor amnii and the fluid of hydatids is similar; se fluid of dropsy is said to contain urea, and to have cholesterine susdin it.
- 46. "Mucus.—This is the secretion of the mucus membranes. When it leaves six or seven per cent. of yellowish solid matter, of which five parts are mucus, the remainder albumen and salts. Mucus does solve in water, but swells like tragacanth into a viscid mass. It distince caustic potash.

17. "Pus is the matter secreted by ulcerated surfaces. When healthy, thick yellowish liquid, formed of opaque globules floating in a clear

When mixed with water, the globules fall, forming a yellow insoludiment. Pus contains about 14 per cent. of solid matter, and is coted by heat and by acids. It contains albuminous matter, fatty matad salts.

18. "The matter of the globules of pus is similar to that of the gloof blood, or globuline. Pus is distinguished from mucus by the miope, or by the action of caustic potash, with which pus becomes thick opy, while mucus forms a thin solution.

### Urine and Urinary Calculi.

- 19. "Urine.—This important excretion is separated from the arterial in the kidneys. It has a pale yellow colour, and a peculiar smell. asity varies from 1.012 to 1.030. It has an acid reaction, or is neutrover alkaline in a state of health.
- io. "On standing, it deposits a slimy mucus-like substance, secreted the lining surface of the bladder. This mucus acts as a ferment, and the urine, after a time, to undergo decomposition; for, when it is uted by the filter, the urine may be kept unchanged for a much longer
- il. "When spontaneous decomposition has taken place, the urine is ne from the presence of carbonate of ammonia, derived from the ureacontains about seven or eight per cent. of solid matter, the remainder water.
- 52. "The characteristic organic principles of urine are urea and uric 5359, 5361). The urea\* has been recently declared by Cap and

fnomalous Cyanate of Ammonia; Urea.—Discovered by Fourcroy and Vauin urine, by Wöhler as the first organic compound artificially produced. It nstituent of uric acid, and is contained in the urine in combination with lactic lenry). Urea is also a product of the reaction of cyanogen on water when a n of that gas is allowed to undergo spontaneous decomposition (Pelouze and deen).

sp.—By mixing fresh urine evaporated to the consistence of a syrup at a gent, which should never reach that of ebullition, when still warm, with its own of colourless nitric acid of sp. gr. = 1.42. If the evaporation has been car-

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Henry to be combined with lactic acid; but, in repeating their exper the editor has always obtained pure urea instead of the lactate. appear, therefore, that in some individuals it occurs uncombined, is

ried sufficiently far, the whole will form a thick crystalline mass; to in small portion of the urine should be tried from time to time. The cryst consists of a compound of nitric acid and urea, which is sparingly solub acid. By the action of the nitric acid on the warm solution, heat is devel affervescence ensues. This is chiefly owing to the destruction of the color ter, and if no external heat is applied, the urea not only is not decor forms, from the first, nearly white crystals of nitrate. When cold is er cording to the method formerly recommended, the crystals are very brown purified with difficulty. It is advisable to separate from the inspisable much as possible of the chlorides it contains, by crystallization, before as mitric sold (Cap and Henry)

"A solution of the colourless crystals of the nitrate of urea is trusted bonats of baryta until it is rendered perfectly neutral; on evaporating, emitrate of baryta, and then of urea, will be obtained. The crystals of the? being rediscolved in a little cold water, are freed from the last portions of \$ of baryta, the solution in alcohol gives crystals of pure urea (Wöhler), states that coloured crystals of urea are best decolorized by a little person potash, which destroys the colouring matter, but has no action on urea.

moon.

"Instead of using nitric acid, the concentrated urine may be added to enturated solution of oxalic acid, when the sparingly soluble oxalits of a which, after being deprived of its colour by charcoal, may be decomposed insoluble oxalits of lime and pure urea, by being digested with pounded the selius). It can also be prepared by the decomposition of the cyanate of silver by sal ammonian, or of the cyanate of oxide of lead by pure or our ammonia."

" Prop.—Crystallizes in colourless, transparent, four-nided, somewhat prisms, of the sp. gr. 1.35, is soluble in its own weight of cold, and in ever tion in hot water, in 4.5 parts of cold, and in 2 parts of boiling alcohol: the solution has a cooling bitter taste like nitre; when pure, it is perfectly a in the air, is not deliquescent, fuses at 250° into a colourless liquid, is deby a higher temperature into ammonia, cyanate of ammonia, and dry mid acid Alkalies do not cause the separation of ammonia in the cold l'a several acids without decomposition to crystallizable saline compounds > rating its solution with nitrate of silver or acetate of lead it is decomposed. ducts being, with the first, nitrate of ammonia and crystalline cyanate of six the second, acetate of ammonia and carbonate of lead. With hyponitron instantly decomposed into nitrogen and carbonic acid gases, which are ev equal volumes, with chlorine it forms hydrochloric acid, nitrogen, and acid. When fused with the hydrated alkalies, or heated in concentrated to acid, it is decomposed together with the constituents of three eq. of water bonic acid and ammonia. Urea contains the elements of evanate of ammonia. + C2NO); it may also be considered, according to Dumas, as a second com carbonic oxide and amide, in which the quantity of the latter is double the amide C<sup>2</sup> O<sup>2</sup> + 2NH<sup>2</sup>

"Nitrate of Urea - This compound, when recently precipitated from a pears in the form of fine crystalline plates of a brown colour and mother lustre, the purer they are, the more they lose this appearance a solution urea treated with nitric acid gives a granular white crystalline precipitate soluble in eight parts of cold, but more freely in hot water, from which it eri in broad, scarcely translucent plates; is sparingly soluble in nitric acid, wit it may be boiled without decomposition. Is composed of one eq of nature a

of urea, and one of water (Regnault)

I am surprised the following process is not mentioned —Impure cyanat ash is prepared by reasting the cyanoferrite of potassium. Aqueous solution cyanate thus obtained, and of sulphate of ainmonia, being mingled, the as is subjected to boiling alcohol, which takes up the urea only On cooling, i crystallizes, and may be rendered purer by recrystallization from the sur struum. Kane, 1164

racid occurs; but, when it is not deposited spontaneously, it appears on the analysis and acid, and the spontaneous deposition of it is probably owing

to the presence of free acid in unusual quantity.

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5853. "The proportion of urea has been found by Lecanu to be tolerably uniform in the same individual, but to vary much in different persons. It is larger in adult men than in women, and least of all in old people and very young children. The proportion of uric acid varies in a similar way. The following analysis of urine, by Berzelius, will give a view of the usual semposition of human urine:—

| Water         |          |      |          |       |         |        |     | 000.00  |
|---------------|----------|------|----------|-------|---------|--------|-----|---------|
|               | •        | •    | -        | -     | -       | -      | -   | 933.00  |
| Urea          |          |      | -        | *     | -       | -      | -   | 30.10   |
| Urie acid     |          |      |          |       | =       | -      | 4   | 1.00    |
| Lactic acid,  |          |      | ammonia, | and   | animal  | mallet | nd- |         |
| hering t      | o them   |      |          | -     | +       | -      |     | 17.14   |
| Mucus of the  | bladd .  | BE   |          | -     |         |        | -   | 0.39    |
| Salphate of   | otaah    |      | -        | -     |         | -      |     | 3.77    |
| Sulphate of   | iods.    |      |          |       | W s     |        | -   | 3.16    |
| Phosphate of  | abon     | -    | -        | -     | -       |        |     | 9.94    |
| Phosphate of  | ammo     | nis  |          |       | -       |        |     | 1.65    |
| Chloride of a | odium    |      |          | -     |         |        |     | 4.45    |
| Hydrochlora   | te of an | nino | tria     | -     |         |        | -   | 1.50    |
| Earthy matte  |          |      |          | pride | of calc | fram.  |     | 1.00    |
| Biliceous ear | th       | -    | -        |       |         | -      | -   | 0.03    |
|               |          |      |          |       |         |        |     |         |
|               |          |      |          |       |         |        |     | 1000.00 |

which gives the colour to inspissated urine, and seems also to be the source of its peculiar odour. He obtained a brown, fusible, resinous mass, having a strong odour of castoreum when dry, and a urinous smell when boiled with water. He calls it oxide of omichmyle, (from anaxes, urine,) and supposes it to contain a radical, omichmyle, the composition of which is still unknown.

5855. "When urine is distilled with an excess of nitric acid, there are formed several products, among which Scharling states that he has observed benzoic acid, and an acid containing chlorine derived from the salts of the urine. This acid appears also to be formed when exide of omichmyle is distilled with nitromuriatic acid. From his analysis, Scharling deduces the formula C<sup>14</sup> H<sup>4</sup> Cl O<sup>2</sup> + HO, which represents benzoic acid, in which one eq. of hydrogen is replaced by one eq. of chlorine. It is also isomeric with chloride of salicule or chlorosaliculic acid (5336).

5856. "Along with this acid there is formed a volatile greenish-yellow oil, which Scharling found to contain twice as much chlorine, and the elements of nitric acid. This compound he calls nitro-chloromichmyle. When bested with acids, it is decomposed, and yields another oily matter, chloro

michmyle. All these observations require confirmation.

5657. "The urine of herbivora is alkaline, and, when the animals are stall-fed, contains, besides urea, hippuric acid; but when they live in the open air, or are forced to labour, benzoic acid alone is found.

5858. "The urine of the carnivora is acid, and contains phosphates and

sulphates of ammonia and soda, as well as uric acid and urea.

5859. "The urine of serpents and of birds is of a soft semisolid consistence, and dries into a mass like chalk. It is almost pure urate of ammonia, but contains a small quantity of phosphates.

acid. They have generally a fawn colour, are soluble in caustic pand and precipitated from the solution by acids. They also dissolve in the acid with the aid of heat; and the solution, when gently evaporated to try ness, leaves a purple stain of murexide. This species of calculus a tody consumed before the blowpipe, leaving a mere trace of ashes.

5861. "Urate of Ammonia occasionally forms a calculus, which we tinguished from the former by giving out ammonia when digested was particularly and particularl

ash.

5862. "Bone-earth forms a common calculus, which is earthy, while in diluted acids, except acetic acid, insoluble in potash, and miestralic

by heat.

5868. "Ammoniaco-magnesian Phosphate also occurs pretty frequent. It is the same double salt which forms whenever magnesia, phosphate acid, and an excess of ammonia, are brought together. It is while a acetic acid, and precipitated again by ammonia. It has often a crystally aspect. When heated, it gives off ammonia, and leaves phosphate of supposia.

5864. "Fusible Calculus.—This very common calculus is a matter of the two preceding. It is white and chalky, and melts easily before to blowpipe. Acetic acid dissolves part of it, hydrochloric acid the rost.

5865. "Oxalate of Lime, or Mulberry Calculus, has a dark-column rough surface, and is very hard. It is insoluble in acetic acid; but when heated to redness, it is converted into carbonate of lime, which dissolute a

acids with effervescence.

5866. "Xanthic Oxide is a rare calculus, first observed by Dr. Mass. It has a light brown colour, and becomes resinous by friction. It denotes in caustic potash, and is precipitated from the solution by carbons and it dissolves in nitric acid without effervescence; and, when evaporate,

leaves a yellow mass. Its formula is C' H' N' O'.

5867. "Cystic Oxide is also very rare. Discovered by Wolfasse. It discovered by Wolfasse. It discovered by Wolfasse. It discovered by Caustic Potash, and is deposited from the solution in hexagonal plates on the addition of acetic acid. It also dissolves in ammonia and the misseal acids; with the latter it forms crystalline compounds. When its solution in potash is heated, ammonia is first given off, and afterwards a combatible vapour, with the odour of sulphuret of carbon. Its formula is C IP NS<sup>3</sup> O<sup>4</sup>.

5868. "Both the preceding species are entirely consumed before the

blowpipe.

5869. "Calculi sometimes occur, in which layers of uric acid alternate with layers of phosphate of lime, ammoniaco-magnesian phosphate, and asible calculus.

#### Changes which occur during the Life, Growth, and Nutrition of Vegetsbles and Animals.

5870. "When we consider that the food of vegetables and of animals is either altogether different from their substance, or passes, before being assimilated, into a new form, we cannot hesitate to admit that the nutrition and growth of both classes of organized beings depend on chemical agencies, although these operate under peculiar conditions, and are inferenced by the unknown force which we call Vitality, so as to produce re-

stalts that cannot be imitated by the chemist in his experiments on dead

5871. "The food of vagetables, as far as their organic structure is sencerned, consists entirely of inorganic compounds; and no organized body can serve for the nutrition of vegetables until it has, by the processes of decay or putrefaction, been resolved into certain inorganic substances.

imown to be the final products of putrefaction. But, even where these are supplied to vegetables, their growth will not proceed unless certain mineral substances are likewise furnished in small quantity, either by the soil, or in the water used to moisten it. Almost every plant, when burned, leaves unless, which commonly contain silica, potash, phosphate of lime; often also magnesia, soda, sulphates, and exide of iron. These mineral bodies appear to be essential to the existence of the vegetable timees, so that plants will not grow in soils destitute of them, however abundantly supplied with carbonic acid, ammonia, and water.

5878. \* In the process of germination, oxygen is absorbed, heat is given out, and in some cases at least an acid, said to be the acetic, is formed, the use of which appears to be to extract from the soil the bases necessary for the future progress of the plant. The starch, or albumen of the seed, becomes soluble, and in the juice undergoes certain changes, by which the woody fibre or lignine, required for the stem and leaves, is produced; but, soon as leaves and roots are developed, the further nutrition of the plant depends on their power of absorbing from the atmosphere and the soil the

matters which constitute the food of the plant.

of the carbon is now derived from carbonic acid, which is either absorbed from the atmosphere and rain-water by the leaves, or from the moisture and air in the soil by the roots. Its carbon is retained, and its oxygen given out; this decomposition being effected in the plant at all times when exposed to the action of light, along with a certain temperature.

5875. "The hydrogen and oxygen of vegetables are derived from water; and the reader will here observe, that the great mass of vegetables, consisting of lignine, starch, gum, &c. is actually composed of carbon

pius water.

from ammonia, which is supplied to them in rain. Liebig has shown beyond all doubt, that rain-water always contains more or less carbonate of ammonia. If we acidulate pure rain-water with a little sulphuric acid, and evaporate to a small bulk, the addition of lime causes the disengagement of ammonia, easily known by its pungent smell. It is remarkable that the ammonia of rain-water has always a putrid smell, which indicates its origin. In fact, it is derived from the putrefaction of preceding races of animals and vegetables, and must at all times exist in the atmosphere; although its relative quantity is so small, that it is not easily detected until it has been accumulated in rain, which, in passing through the air, dissolves it readily, and conveys it to the earth.

6677. "It is also to be observed, that the soil itself, like all porous bodies, possesses the property of absorbing ammonia, and therefore will attract it from the atmosphere. Alumina, peroxide of iron, and humus, all absorb ammonia powerfully. Gypsum (sulphate of lime) and other sulphates convert the carbonate of ammonia into the more fixed sulphate, which remains

in the soil till absorbed by the roots. This explains in a great me

use of these ingredients in fertile soils.

5878. "It is only under the influence of light that plants can desspose carbonic acid, fixing its carbon and setting free its oxygen. Dung the night, on the contrary, they undergo a kind of slow combustion, or the being absorbed, and carbonic acid formed. But the halance in this current alternation is vastly in favour of the process by which oxygen a sext and the atmosphere, for, the whole carbon of a forest, for example, being anvel from carbonic acid, an equivalent quantity of oxygen must have been liberated; and this consideration alone enables us to explain the fact, and, notwithstanding the enormous amount of oxygen withdrawn from the 🛸 mosphere by the respiration of animals, by combustion, by putrefactors, and by the action of vegetables during the night, in all of which process the oxygen is converted into carbonic acid of equal volume, the proported of oxygen in the atmosphere does not diminish, and that of carbonic will does not increase.

5879. "From these considerations it appears that there must always exist a balance or fixed proportion between the existing amount of animi and that of vegetable life. Where animals abound, and where men cary on the usual operations of civilized life, there, carbonic acid must be larger formed. But this carbonic acid, in yielding its carbon to vegetation, page also its oxygen to restore the purity of the air, and support again the ration of mea and animals. Again, the decay and putrefaction of in the mals and vegetables yield carbonic acid and ammonia, the very subserve which form the food of a new race of vegetables; and these again bute to the nourishment of new animals; so that, in this uncessary of chemical changes, the death of one generation supplies the mean of in

to that which is to follow.

5880. "It has long been the prevailing opinion, that the carbon of place is derived directly from humus or humic acid existing in the soil, which # supposed to be absorbed in the form of a solution in water, or as human of ammonia; but it must be admitted, as Lieb g has shown, that there a 🚥 evidence whatever that humus is directly absorbed by plants. Hims. \*\* it exists in the soil, is almost entirely insoluble in water, and, when a sourble form occurs, the solution, however weak, is always of a dark brown colour; whereas the juices of plants, when first absorbed, are coordinate Again, humic acid, as described by chemists, never occurs in sods, but #1 product of the action of alkalies on humus, and besides forms solubles = dark-coloured as those of humus. Good fertile soil digested with cod #15 ter yields to it no colour; water, filtering through such soil, passes cooks less, as may be daily observed; nay, moss-water, which is actually coloured brown by humus, is decolorized by passing through a good feruic 🗪 containing humus; finally, a peaty soil, which contains more humes 200 any other, is notoriously barren.

5881. "On the other hand, the first vegetables which grew on the earth could not have derived their carbon from humus, which is a product of the decay of vegetables, but could only have obtained it from carbonic and and if this source of earbon were then sufficient, there is no reason to leek for another. Besides, if we reflect on the extreme luxuriance of regential in uninhabited countries, where the soil has never been manufed, we cannot fail to perceive that the carbon of that vegetation must have been theft derived from the atmosphere; and when, in addition to this, we had the the proportion of humus in all soils bearing vegetation increases rather that

diminishes, in spite of the vast amount of carbon annually accumulated and removed in the crops, we are compelled to adopt the same conclusion.

5882. "This latter consideration shows, that the humus and other orpatic matters in manures do not act directly in furnishing carbon, and that their use chiefly depends on other ingredients. These, as Liebig has dethoustrated, are, first, the ammonia they contain or yield by putrefaction; and, secondly, the mineral bodies, such as potash, phosphate of lime, &c., found in their ashes.

5883. "But, although there is no evidence that humus is directly absorbed by plants, and the phenomena of peat and mossy soils prove, that the poluble forms of humus are unfavourable to vegetation, yet it cannot be toubted that humus or mould, both of the soil and the manures, performs an important function. It slowly and gradually undergoes combustion, yielding a constant and steady supply of carbonic acid in moderate quantity. This is partly absorbed by the roots, and partly rises into the atmosphere to be absorbed by the leaves; but, as the proportion of humus in the toil does not diminish, that which is thus consumed is probably restored to be soil by the secretions, or rather excretions, from the roots.

5884. "Humus also probably acts by absorbing and fixing the ammonia

of the atmosphere.

5885. "According to the views above stated, which have been admirably laid down by Liebig in his Agricultural Chemistry, the chief use of natures is not to supply plants with carbon, but with ammonia and inormaters. Every plant requires certain mineral substances, without which it cannot prosper; and a soil is fertile or barren for any given plant, according as it contains these. Thus, the ashes of wheat-straw contain such silica and potash, while the ashes of the seeds contain phosphate of ammonia and magnesia. Hence, if a soil be deficient in any one of these, t will not yield wheat. On the other hand, a good crop of wheat will exsaust the soil of these substances, and it will not yield a second crop till hey have been restored, either by manure or by the gradual action of the reather in disintegrating the subsoil. Hence the benefit derived from fallows and from the rotation of crops.

5886. "When, by an extraordinary supply of any one mineral ingrelient, or of ammonia, a large crop has been obtained, it is not to be expected that a repetition of the same individual manure next year will produce he same effect. It must be remembered, that the unusual crop has exmusted the soil probably of all the other mineral ingredients, and that they

and must be restored before a second crop can be obtained.

**5887.** "The salt most essential to the growth of the potato is the double **bosphate** of ammonia and magnesia; that chiefly required for hay is phoshate of lime; while for almost all plants potash and ammonia are highly expericial.

5888. "From the principles above mentioned we may deduce a few raluable conclusions in regard to the chemistry of agriculture. First. By reamining the ashes of a thriving plant, we discover the mineral ingredients which must exist in a soil to render it fertile for that plant. Secondly. By reamining a soil, we can say at once whether it is fertile in regard to any slants, the ashes of which have been examined. Thirdly. When we know he defects of a soil, the deficient matters may be easily obtained and added o it, unmixed with such as are not required. Fourthly. The straw, caves, &cc. of any plant must be the best manure for that plant, since every regetable extracts from the soil such matters alone as are essential to it.

This important principle has been amply verified by the success standing the use of wheat-straw or its ashes as manure for wheat, and of the depings of the vines as manure for the vineyard. Where these are use, as other manure is required. Fifthly. In the rotation of crops, those shall be made to follow which require different minerals; or a crop where tracts little or no mineral matter, such as peas, should come after use what

exhausts the soil of its phosphates and potash.

the phosphates, which have been extracted by successive crops of gas and corn, the whole of the bones of the cattle fed on these crops having the rived from the soil; its gelatine also yields ammonia by putrefacture. Co ano acts as a source of ammonia, containing much oxalate and unit of ammonia with some phosphates. Night-soil and urine, especially the ter, are most valuable for the ammonia they yield, as well as for phosphate and potash: but are very much neglected in this country, although the importance is fully appreciated in Belgium and China. Brun is a very valuable manure, especially for potatoes, as it contains much of the ammoniaco-magnesian phosphate.

5890. "Nitrate of Soda probably acts by its alkali, replacing pound but it is possible that its acid may also yield nitrogen to plants, aking we possess at present no evidence of this, and indeed no evidence be plants can derive their nitrogen from any other source than from any

5891. "Such is a brief sketch of the general laws of vegetator at a present known, in so far as they are connected with chemistry. Of the changes in the juices of vegetables, by which the numerous products of the vegetable kingdom are formed, we know nothing. The juices of place contain ammonia and sugar, gum or starch; all the elements are therein present from which the nitrogenized compounds, albumen, fibrus, and caseine, in other words, proteine, may be formed, and it appears that the tables alone can produce proteine. Thus the final products of regular form the food of animals; the modifications of proteine serving for maniform the food of animals; the modifications of proteine serving for the support of respiration.

5892. "The life of animals is distinguished chemically from that of vegetables by the circumstance, that in the former oxygen is company absorbed and replaced by carbonic acid, while in the latter carbonic acid absorbed, its carbon retained, and its oxygen given out. Consciousness and

the power of locomotion are peculiar to animals.

5893. "In animals two processes are constantly carried on; that of respiration, by which the animal heat is kept up, and that of nutrition, by which the matter consumed in the vital functions and expelled from the

body is restored.

5894. "Respiration is essentially a combustion of carbon, which is combining with oxygen is converted into carbonic acid, and at the same the furnishes the animal heat. Liebig calculates that the amount of carbon daily burned in the body of an adult man is about fourteen ounces, and that the heat given out is fully sufficient to keep up the temperature of the heat, and to account for the evaporation of all the gaseous matter and water expelled from the lungs.

5895. "This carbon is derived in the first place from the tissues of the body, which undergo a constant waste, but ultimately from the field.

5896. "In the carnivora, whose food is almost entirely composed of compounds of proteine, albumen, &c. one part is devoted to supply to

waste of the tiesues, while another portion, or a corresponding amount of previously existing tissue, is decomposed so as to yield the carbon required for respiration. As the tissues can only be decomposed by the exercise if the vital functions, this is the reason why, in the carnivors, an enorthous amount of muscular motion is required to furnish the necessary supby of carbon-

5897. "On the other hand, the food of the herbivora contains but little If the compounds of proteine, only sufficient to restore the waste of the houses; while the carbon required for respiration is supplied by the starch, men, sugar, oil, &c. which form the great mass of their food, and no such smount of muscular motion is required in them as in the carnivors.

5898. "It is in the form of bile chiefly that the carbon undergoes commation. Hitherto, the true function of the bile has been disputed; and by agest authors that fluid has been considered as an excretion, intended to be appelled from the body in the faces. But Liebig has shown that only a mall fraction of the whole amount of bile can be detected in any shape in in freeces, and that the bile unquestionably is resheorbed in the intestinal amai, and re-enters the circulation, where it soon disappears; and as the reportion of carbon in the bile is very large, although not sufficient to account for all the carbonic acid given out, there is no reason to doubt that is gradually consumed by the oxygen of the arterial blood, and convert-I into carbonic acid and water, which escape by the lungs and skin.

\$899. "To return to the subject of the animal heat: the food that is equired, and hence the appetite, must be proportional to the amount of arbon required to supply the animal heat. Now, in hot climates, where he external cooling is less, less heat is required, the appetite is much more soble, and the usual food, consisting of fruits and vegetables, contains a ar smaller amount of carbon than in cold climates, where the appetite is men, and the food highly carbonized, such as flesh, or even blubber. For be same reasons, warm clothing, by diminishing the loss of heat by exarmal cooling, blunts the appetite; and those who remove from a cold to a were climate always find that their appetite fails. This is a warning from ature to diminish the amount of food taken; and if it were attended to, and the common but absurd practice of stimulating the appetite by ardent quore and hot spices abandoned, Europeans might enjoy as good health a the East or West Indies as at home. It is obvious that, even in Europe, sore food is required in winter than in summer. (Animal Chemistry, 28.) 5900. "In endeavouring to explain the formation of the bile, it is obinusly of no moment whether we derive it from the albumen, fibrine, dec. f the food, or those of the tissues, their composition being identical. Liebig, seaming choleic acid to be the chief organic constituent of the bile, and its nemula to be C76 No Hea Ose, has shown that the half of this formula, added > that of urate of ammonia, C<sup>10</sup>H<sup>2</sup>N<sup>7</sup>O<sup>6</sup>, which given the sum C<sup>40</sup>N<sup>6</sup> [40 O<sup>17</sup>, is equal to the formula of blood or flesh, C<sup>40</sup> N<sup>4</sup> H<sup>20</sup> O<sup>15</sup>, with the adition of one atom water and one atom oxygen. (Animal Chemistry, 185, 86.) Again, proteine, C40 No Had O14, plas three atoms of water, gives 30 same sum, excepting one atom of hydrogen, viz. C40 N5 H40 O17. is way we can see how the tissues, acted on by oxygen and water, may juid the ingredients of bile and urine. This is the first attempt which has con made to trace chemically the connection between the food, the blood r the tissues, and the secretions or excretions; and showing, as it does, ant these questions are capable of elucidation on chemical principles, it

must be regarded as the most important idea yet suggested in animal de-

mistry.

5901. "Supposing it to be well-founded, the tissues which are considered resolved first into bile and urate of ammonia. The former is secret from the liver, reabsorbed and burned, as before stated. The latter, a sepents and birds, is expelled unchanged; but in man and quadrooms whom the amount of oxygen inspired is much greater, it is also united.

yielding finally carbonic acid, ammonia, and urea.

5902. "Should the supply of oxygen in the human subject to make cient to act on the urate of ammonia, then the tric acid is deposit to gravel or calculus; if the supply of oxygen be somewhat greater, but all deficient, oxalic acid is the result, and mulberry calculus occurs; but a much exercise be taken and abundance of oxygen supplied, the output of the uric acid is completed, and nothing is left but uren or carbona ammonia.

5903. "This explains the true cause of uric acid and mulberry calculate to be a deficiency of oxygen; it also explains why uric acid calculate is lowed by mulberry calculus in those who remove from the town to be country, where more exercise is taken; and from these considerations we may see how valuable are the results which will flow from a thorough or

vastigation of all departments of animal chemistry.

ing, besides urea, hippuric acid when they are at rest or stall-fed, and sozoic acid when they are in full exercise, and when consequently were
oxygen is supplied. Liebig has shown, that, if to five times the formal
blood, we add nine atoms of oxygen, we have the elements of an accompanie, and three atoms water; and that, if to five times the formal a
blood we add forty-five atoms oxygen, we obtain the elements of an accompanie acid, thirteen and a half atoms urea, three atoms cholenc acid, him
atoms carbonic acid, and twelve of water. Moreover, two atoms process
with two atoms of water, contain the elements of six atoms allentime to the urine of the fætal calf), and one atom cholondic acid, which a sepposed to be the same as the meconium.

the carnivora, an ox secreting, according to Bardach, 371bs, of but data. As the waste of matter in the herbivora is but limited, it is obvious that cannot supply all the bile, and consequently a great part of it must be rived from the starch and other nonazotized constituents of their food, what lose oxygen, and enter into combination with some azotised product of the product o

decomposition of compounds of proteine.

5906. "In order to show how this is possible. Liebig points out that the elements of two atoms of proteine, with those of three atoms une and is two atoms oxygen, amount to the same sum as six atoms hippure and and nine atoms urea; while, if to five atoms starch we add two atoms impure acid and two atoms oxygen, the sum is equal to two atoms choice and twenty atoms carbonic acid.

being present, undergo transformation, and mutually affect each other, or products of this metamorphosis may be urea, choice acid, ammona, as

carbonic acid. Thus:

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5 atoms proteine, 5 (Ca Ne Hat O14) = Cate Nat Hate O70
                      starch, 15 (C^{12} H^{10} O^{10}) = C^{130}
                                                               H150 O150
            15
             12
                                                               His Ois
                      water, 12 (
                                           H O =
                      oxygen,
                     The sum is
                                                   - C400 Nao Hasa O237.
* And
                                       9 (C \approx N H \approx O = C \approx N \approx H \approx O \approx
          9 atoms choleic acid,
                                       9 (Cs Ns H4 Os) = Cis Nis Hst Ois
                   urea,
                                              N H
                   ammonia,
                                                      O^2) ==
                   carbonic acid,
                                      60 (C
                                                                           Oim
               The sum is
                                                          Con Nao Hats Ocal
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5008. "The reader will observe that these equations are given, not as representing what is actually proved to occur, but only to show how such changes may be conceived on ordinary chemical principles. But it is to be porme in mind, that all the necessary substances 'meet in the circulation; procesine and starch from the food, oxygen in the arterial blood, and that water is never absent: while the resulting products are the chief constituents of the secretions and excretions; viz. carbonic acid, excreted by the lungs; zrea and carbonate of ammonia, excreted by the kidneys; and choleic acid, perceted by the liver. (Animal Chemistry, 150 et seq.)

5909. "We have thus seen how the carbon in the form of choleic acid bile, may be obtained in a state most favourable for its oxidation or combeastion. But, if the supply of oxygen be deficient, the choleic acid may by a partial oxidation yield hippuric and lithofellic acids, just as we have seen that uric acid, partially oxidized, yields oxalic acid; for two atoms choleic acid + 10 atoms oxygen are equal to two atoms hippuric acid, one atom lithosellic acid, and sourteen atoms water. Thus one species of biliary calculus, identical with the bezoar stones found in the herbivora, may have an origin similar to that of the mulberry calculus, both arising from a deficient supply of oxygen. (Liebig.)

5910. "Soda is necessary to the formation of bile, and is supplied in the form of common salt. Where the supply of soda is defective, the metamorphosis of proteine can yield only fat and urea. If we assume for fat the empirical formula C11 H10 O, then two atoms proteine, with twelve atoms water, and fourteen atoms oxygen, in all Cos Nas Hot Ost, are equal to six atoms urea, six atoms fat, and eighteen atoms carbonic acid. If we assume fat to be C19 H10 O, a similar result may be traced; and the composition of all fats lies between these two empirical formulæ. Now, it is worthy of obpervation, that, if we wish to fatten an animal, we must carefully avoid

giving much salt in its food. (Liebig.)

5911. " As another point of connection between the products of the metamorphosis of bile and of the constituents of urine, in addition to the possibility already mentioned of both being derived from the oxidation of proteine, it may here be remarked, that three atoms taurine and three atoms ammonia are equal to one atom uric acid, one atom urea, and twenty-two atoms water; and that one atom taurine and one atom ammonia are equal to one atom allantoine and seven atoms water. (Liebig.)

5912. "It may further be noted that one atom uric acid, fourteen atoms water, and two atoms oxygen, correspond to two atoms taurine and one atom urea; or, if two atoms water be added, to two atoms taurine and two atoms carbonate of ammonia. Moreover, one atom alloxan and ten atoms water are equal to two atoms taurine; and one atom taurine contains to elements of two atoms oxalic acid, one atom ammonia, and four atom ter. (Liebig.)

5913. "As allowed is a product of the exidation of uric acid, and as has been shown above to be related to taurine, that is, to bile, it work to very important to study its action on the system. It might probably as beneficially in some discusses of the liver. It may be suffely adminished a

considerable quantity. (Liebig.)

5914. "In the urine of the carnivora we find soda in moderate quarty, combined with sulphuric and phosphoric acids. This soda was commit in their food, and, after contributing to form the bile, has been secree to the kidneys. But it is never sufficient to neutralize the acids protect and consequently we find much ammonia along with it, while the same acid.

5915. "But in the urine of the herbivora soda is present in far high quantity, and combined with carbonic, hippuric, or benzoic acd. The shows that the herbivora require a far greater amount of soda than a catained in the amount of blood daily consumed, which in them is small, at this soda is obtained from their food, and employed in producing that the dant bile.

destitute of alkalies; but these alkalies are not less necessary for the seport of the animals than of the plants. The soda is found in the bind sole; and the potash is now known to be absolutely essential to the praction of caseine, that is, the secretion of milk. In like manner, the phophet of lime, which is essential to the growth of grasses, as equally cases the production of bone in the animals which feed on these plants. It are possible not to be penetrated with admiration of the wisdom which is shown in these beautiful arrangements.

5917. "Let us now consider the changes which the food undergood the process of digestion, and we shall observe this process in the carriers where it is most simple, as their food is identical in composition with the

tissues.

poured out, and after a short time the whole is converted into a sending homogeneous mass, the chyme. Many researches have been made to decover the solvent contained in the gastric juice, but in vain. It contains the substance which has the property of dissolving fibrine, albumen. Acc., are we are compelled to adopt the opinion of Liebig, according to which the food is dissolved in consequence of a metamorphosis, analogous to fermination, by which a new arrangement of the particles is effected. As, a fermentation, the change is owing to the presence of a body in a same of decomposition, or motion, which is propagated from the ferment to the sager by contact; so, in digestion, the gastric juice contains a small quantity of a matter derived from the lining membrane of the stomach, which is in a case of progressive change, and the change or motion is propagated from the to the particles of the food under certain conditions, such as a certain temperature, &c.

5919. "The phenomena of artificial digestion confirm this view. If the lining membrane of a stomach, perfectly clean and fresh, be infined in the ter feebly acidulated with chlorohydric acid, the liquid acquires no notices action on albumen; but if the membrane be exposed to the air for some time, or be left in water for a while,—in short, if decomposition be allowed

commence,—then the infusion, if coagulated albumen or fibrine be placed it, and the whole kept at the temperature of the body, by degrees effects

perfect solution or digestion.

5920. "Prout has shown that the gastric juice contains free chlorohyic acid. This is derived from the common salt, the soda of which comness with the albumen or fibrine, while its acid, being set free, at length
r its accumulation checks further change. Besides the gastric juice, the
ly other substance employed in digestion is the oxygen which is introneed into the stomach with the saliva, which from its viscidity encloses a
rge quantity of air. The chyme then leaves the stomach, and gradually
ness into the state of chyle, which resembles blood, except in colour, being
leady alkaline, not acid like the chyme.

5921. "By means of the circulation oxygen is conveyed in the arterial bood to every part of the body. This oxygen, acting on the tissues descended to undergo change, produces a metamorphosis, by which new soluble empounds are formed. The tissues thus destroyed are replaced by the matter derived from the food. Meantime, those of the products of memorphosis which contain the principal part of the carbon, are separated om the venous blood in the liver, and yield the bile; while the nitrogen ocumulates, and is separated from the arterial blood in the kidneys in the

mm of urea or uric acid.

5922. "It has been already mentioned, that vegetables alone possess the ower of forming proteine, which they furnish to animals in the forms of burnen, fibrine, and caseine. In the animal body these forms of proteine re employed to yield the different tissues, most of which bear a simple retion to proteine. Thus,—

Fibrine, albumen, caseine, are

Arterial membrane is

Chondrine is

Pr + S + P + salts.

Pr + 2HO.

Pr + 4HO + O<sup>2</sup>.

Pr + 4HO + O<sup>2</sup>.

Pr + NH<sup>3</sup> + O<sup>2</sup>.

Gelatinous tissues are

Pr + S + P + salts.

Pr + 2HO.

Pr + 4HO + O<sup>2</sup>.

Pr + NH<sup>3</sup> + O<sup>2</sup>.

5923. "It is not meant that these formulæ express the actual constitution f the tissues, but only that they give the proportion of the elements actually resent, and show how they might give rise to the tissues. Some of these issues contain proteine, or at least yield it when acted on by potash: this the case with hair and horn. But others, as, for example, the gelatinous issues, although doubtless derived from proteine, do not contain it, and consquently cannot yield any of its modifications. This explains the fact hat gelatine alone cannot support animal life. It cannot yield blood or suscular fibre, although it may serve to nourish the gelatinous tissues. See Gelatine.)

5924. "Liebig has shown (Animal Chemistry, p. 141), that gelatine may be formed from proteine in two ways; either by adding to two atoms proteine three atoms allantoine and three atoms water (which are equal to me atom uric acid, one atom urea, and four of water), or by subtracting from three atoms proteine half an atom choloidic acid, and adding four stoms water. These statements apply to Mulder's formula for gelatine; but as the true formula is still doubtful, they are only mentioned to show the method by which we may hope to arrive at accurate results.

5925. "There is another constituent of the animal body, namely fat, the production of which deserves notice. It is not an organized tissue, but is formed and collected in the cellular tissue under certain circumstances.

These are rest and confinement, that is, a deficiency of oxygen, and a abundance of food devoid of nitrogen. Carnivorous animals are new to

and the herbivora only become so in confinement.

of which is such, that, if deprived of oxygen, fat remains. If from and C<sup>15</sup> H<sup>10</sup> O<sup>10</sup>, we take nine atoms oxygen, there remains C<sup>15</sup> H<sup>10</sup> U with is one of the empirical formulæ for fat. Or if from starch we remainder O<sup>15</sup> H<sup>10</sup> U with atom carbonic acid, CO<sup>0</sup>, and seven atoms oxygen, the remainder O<sup>15</sup> H<sup>10</sup> U represents the other empirical formula of fat. We have already so have fat may be derived from proteine when sode is deficient; and we may have add, that all the elements of food contain more oxygen than fat, in propertion to the carbon. Thus, in albumen, fibrine, and caseine, for 120 a carbon there are contained 86 eq. oxygen; in starch, for 120 eq. oxygen; and in grape-sugar, 140 eq. oxygen; while in fat there are only 16 a oxygen for 120 eq. carbon.

of deoxidation. But we have seen that it is produced where oxygen and ficient; and it appears, as Liebig has pointed out, that when there is a strict supply of oxygen, the production of fat, which is the consequent of this deficiency, yields a supply of that element, and thus serves to imput the animal heat and the vital functions, which would otherwise be according to this is another beautiful instance of contrivance, equally sample at well.

derful-

5928. "That fat must be formed by the deoxidizing process about bluded to, is proved by the phenomena of the fattening of ansats. A goose, tied up, and fed with farinaceous food altogether destitute of it, to quires, in a short time, an increase in weight of several pounds, the wish of which is fat. Again, the bee produces wax, a species of fat, from pure

sugar.

alone can form, we see, from its composition, intermediate between the proteine and fat, that it may be formed either by depriving protein a some nitrogenated product, or by adding such a product to fat. When it is formed we do not know; but it must be formed in the animal body: the Liebig has suggested that the power of the vegetable alkalies to affect to nervous system, may be owing to their composition, which approach nearer to that of nervous matter than any other compounds. There are lies may promote or check the formation of nervous matter, and thus produce their peculiar effects.

5930. "In like manner, certain vegetable products analogous to the egetable alkalies, such as caffeine (or the ne) and the observance, may be see posed, according to Liebig, to promote the secretion of bile, their components.

sition being related to that of some of the products of bigs.

bromine, with twenty-two atoms water and sixteen atoms oxygen, over ponds to four atoms taurine and one atoms oxygen, over eight atoms water, and fourteen atoms oxygen, contain the elements of the atoms taurine and one atom oxygen, contain the elements of the atoms taurine and one atom unce eight. (Atoms the elements of the atoms taurine and one atom unce acid. (Atoms Chemistry, 150.

5932. "Now it is surely very remarkable, that the vegetables contains these compounds, tea, coffee, and cocon, should be, one or other of these used by almost all nations to yield a refreshing drunk; and it is so, as the compounds."

mericus that the peculiar principle of tea should turn out to be identical with

that of coffee, as recent researches have demonstrated.

1933. "We may suppose, with some degree of probability, that where the formation of bile, and consequently that of urine, which is connected with it, does not go on as it ought, the use of these beverages, by promoting the secretion of bile, may assist the process of respiration, promoting the mirrial heat, and, at the same time, contributing to the due performance of the vital functions. At all events, neither the beneficial and refreshing tificats of these articles of diet, nor their relation to bile and urine, can be pre-rlooked; and the universal adoption of the practice of using tea, coffee, or chocolate, is a proof that men have discovered and obtained from different sources the means of producing the same effect.

5934. "The preceding observations are sufficient to show that we may expect, in progress of time, to explain the action of all remedies on chemical principles. The true path has been opened up, and it only remains for ex-

perimenters to pursue it with energy and perseverance."

#### OF RESPIRATION.

5935. The quotation from Gregory's work being concluded, I subjoin an article which I had prepared on respiration, as it contains some ideas which are not found in the preceding matter, and some objections to Liebig's ex-

planation of the phenomena of that process.

the volume of the air respired by animals is diminished, but that a portion of the oxygen is replaced by an equal bulk of carbonic acid. Although, at one time, by respectable observers, the volume of this last mentioned gas was alleged not to be uniformly equal to that of the absorbed oxygen, the ratio of the one to the other, being represented as varying with the time of day and the season, not only in different animals, but also in the same animal, later observation seems to have produced a general opinion, which is zealously espoused by the distinguished chemist above mentioned, that the expired carbonic acid is, upon the whole, exactly equivalent to the oxygen consumed.

5937. The prevalence of nitrogen, in animal substances, naturally led to the idea that it might be assimilated more or less during respiration; but experience has led to an opposite opinion; and Liebig has endeavoured to show, that in the nutriment of granivorous animals, there is no deficiency of vegeto-animal matter having as large a pro-

portion of nitrogen as flesh and blood\* (5023).

I subjoin the following opinion of Berzelius. Report for 1840, page 313.
 The question has often been put, whether animals assimilate nitrogen during respiration. In examining air which has been breathed by them, it has been found



5938. When first, by the Lavoiserian school, the best of all ordinary fires was shown to be attributable to the union of oxygen with the combustible employed, the dest naturally followed, that respiration being attended by a like union of oxygen with combustible matter, animal best ought to be ascribed to this source. Many objections to this explanation of the origin of animal heat were subsequently urged, and, among others, the fact that the best of the lungs, the fire place, is no higher than remoter parts of the animal frame.

5939. To remove this objection, Crawford suggests that the capacity for heat, of arterial blood, being greater than that of venous blood, caloric was taken up by the blood in one state, to be evolved when in the other. The suggestion respecting the relative capacities for heat, of arterial and venous blood, has not been supported by subsequent experience; and another view of the subject has been taken, which renders it quite consistent that the perature should not be peculiarly high in the lungs.

5940. It is supposed that the blood merely absorbs any gen in the lungs, but that this oxygen is carbonized durate its circulation, and thus causes heat to be given out in a parts of the system. The carbonic acid thus produced, a reaching the lungs in combination with the venous blook is exchanged for oxygen, and consequently expired with

the breath.

of the red globules is held by the arterial blood, in the state of hydrated sesquioxide; but in the capillaries, the sesquioxide passing to the state of protoxide, by yielder oxygen to the carbon in the blood, combines with the car

that in some cases a deficit of nitrogen has ensued, in others an excess, which others, again, the proportion has remained unchanged. Let rigorous expension have proved, that the nitrogen of respired air is quite passive, and cannot be small lated during respiration. Increaver, that the blood in common with all other bytes in contact with the air, contains nitrogen and oxygen in the proportion is they are present in the gaseous mixture employed so that when a mixture taining more nitrogen, is respired, a greater quantity is absorbed. When the with ture, under like circumstances, has an inferent quantity of nitrogen, this property given out by the blood. It may be assumed, that experiments have competed acided that the proportion of nitrogen in the animal frame is altogether independent of the quantity of air respired."

It does not, however, appear to me to be true—that all liquids in contact with we take up its ingredients in the same proportion, or if they do, that they contains hold them in that proportion, uninfluenced by the chemical affinity between the

constituents and oxygen

bonic acid thus produced, and gives rise, in the venous

blood, to a carbonated protoxide.

5942. When the venous blood reaches the lungs, the protoxide exchanging carbonic acid for oxygen, this gas is expelled with the breath, while the regenerated sesquioxide again, by union with water, reconverted into a hydrate. The well known change of hue which follows the transfer of the blood from the veins to the arteries, through the pulmonary organs, seems to be considered as a collateral consequence of these chemical reactions. Yet this change does not appear to me sufficiently accounted for, since no such alteration of colour can be produced by the transformation of a carbonated protoxide of iron to a hydrated sesquioxide. Moreover, the fact that no peculiar elevation of temperature takes place on the surfaces where the venous blood meets the breath, seems to me inconsistent with Liebig's explanation, since the heat must be extricated in the space where the iron is peroxidized.

5943. Upon the whole I now think as I have for forty years, whatever other opinions may have prevailed, that there must be a degree of heat derived from respiration proportioned to the quantity of oxygen converted into carbonic acid; but with all due deference for Liebig, I do not agree with him, that it is possible to give a satisfactory explanation of this process upon purely chemical affinities, such as exist independently of vital power. It appears to me that nature has the power, within certain limits, of making chemical affinities to suit her own purposes, and can therefore cause the oxygen to be absorbed, the carbon to combine therewith, and the heat to be given out when and where the processes of vitality require it. If nature have not the alleged power, how does it happen that, out of the heterogeneous congeries of elements existing in the egg, the bill, the claws, the feathers, the bones, the blood, and flesh, are made to appear at the various stations, at which their presence is requisite, for the existence of a young bird?\*

<sup>•</sup> Mr. Winn, (L. and E. Phil. Mag. 174,) considers the extension and contraction of the fibrous tissues of the arteries, during pulsation, as among the causes of animal heat. It is well known that caoutchouc grows warm when rapidly extended; and Mr. Winn found a portion of the aorta of an ox to be capable of a similar rise of tem. perature, when, during two minutes, it was made to undergo turgescence, and col. lapse similar to that which takes place during pulsation. To have decided this ques-

5944. Liebig cites the following interesting facts. active man expires 13.9 ounces of carbon, and daily sumes, in the same time, 37 ounces of oxygen = 51 cubic inches, or about 223 gallons. Reckoning 18 is rations per minute, there must be 25,920 consumed day, and consequently \frac{11448}{11448} = 1.99, or nearly two consinches of oxygen in each respiration. In one minute therefore, there are added to the blood 1.99 × 18 = 5 cubic inches of oxygen, weighing rather less than two grains.

5945. In one minute, ten pounds of blood pass throthe lungs, measuring 320 cubic inches, among which being divided, there must be one cubic inch of oxygen

nine of blood nearly.\*

5946. Ten Hessian pounds of blood = 76,800 grains the arterial state, contain 61% grains sesquionition; if in the venous state, 55% of protoxide of 6% difference, is the quantity of oxygen which the iron of venous blood can acquire in the lungs, which, definition twelve grains, the whole quantity of oxygen about leaves 5.60 grains requiring some other means of about tion. But 55% grains of protoxide of iron would take 73 cubic inches of carbonic acid, which is double the lume that the 35% of oxygen can generate.

5947. One glaring defect in this part of the explanations from the admitted fact, that nearly one-half of absorption of oxygen is unaccounted for; 5.60 in two

parts.

#### OF FERMENTATION.

5948. Certain spontaneous changes which ensue in organic subst by which they are more or less decomposed or resolved into new contions, have been generically designated under the name of fermentati

5949. For a long time only three kinds of fermentation had been nised, called, severally, the vinous, the acetous, and the putrefactive now we have several others added to the list, among which are the serine, and the viscous or lactic.

5950. The production of cyanhydric acid (1323) by the reacti

tion, the author should have shown that heat might be permanently caused extension and contraction either of caoutchouc or the ox artery. But were monstrated that heat could be thus permanently generated, there would be t difficulty in explaining how the organic substances employed could thus give heat. It involves the question of the materiality of caloric, since, if material, manent supply could not be derived from an isolated strip of caoutchouc.

<sup>\*</sup> Stated upon the authority of Muller. "Physiologie, Vol. 1, p. 345." † Deduced from the Researches of Dénis Richardson and Nasse, Handworte der Physiologie, Vol. 1, p. 138. Note.—Measures and weights are Hessian

sulsine with amydaline (3055), that of the oil of mustard by myrozine nd myronic acid (5091), are, by Boutron and Fremy, considered as cases fermentation; and to these, it seems, we may add the generation of nitin, which is alleged to be the effect of a species of fermentation promoted the leaf of the tobacco plant after it has been gathered.

5951. To the saccharine, the vinous, acetous, and viscous or lactic ferentation, allusion has already been made in treating of starch (4082); of ne sugar (4057); of alcohol (5578); of lactin (4070); acetic acid 197); lactic acid (5215).

# Of the Saccharine and Vinous Fermentations.

5952. The saccharine fermentation is exemplified in the change which kes place in the mash or wash of the distiller, by which the starch of the ain, C19 H10 O10, takes two atoms of water, 2HO, to form dry grape su-17, C10 H19 O19.

5953. The vinous fermentation ensues in all cases where alcohol is proseed by an internal change in organic solutions. By some chemists, it is pposed that alcohol is produced only when grape sugar is present at the stact, or generated subsequently; since it is alleged that cane sugar and her saccharine substances must be converted into grape sugar before they n enter into the vinous fermentation. It has been stated, that by this rmentation an atom of grape sugar is resolved into the elements of two oms of alcohol and four atoms of carbonic acid (5578).

5954. The juice of the apple, the pear, or the grape, at any temperature pove 50°, spontaneously enter into the alcoholic fermentation. cribed to the existence, in them, of a vegeto-animal matter, which being est oxidized, afterwards mysteriously causes the sugar to be resolved into cohol and carbonic acid, as already stated (5578). The preservation of uits and other organic substances by heat, in well closed vessels, is asibed to the prevention of that oxidizement of the vegeto-animal ferment,

bich is the necessary precursor of fermentation.
5955. In the case of wort as prepared in breweries, there is great diffiilty in inciting a proper vinous fermentation, without the assistance of east arising from a preceding process. Yet during every well conducted peration, a large quantity of this substance has to be thrown off. The orough performance of this process, called cleansing, has always been nown to be necessary to the flavour of the beer; but Liebig alleges that it so lessens the liability to acetification, and that by a process practised in avaria, the yeast being more thoroughly removed by deposition, such a periority was attained as respects insusceptibility of sourness, that large emiums were offered in other German states for those who should suced in imitating that process. This consists in the exposure of the beer, in en shallow vessels, to atmospheric oxygen, at a temperature below 50°, which the vegeto-animal matter which forms the yeast, is oxidized and ecipitated at a temperature too low for that simultaneous conversion of the cohol into acetic acid, which would be the consequence of a higher temrature under like circumstances.

5956. During fermentation there is a commensurate attenuation of the quor, of which the extent may be ascertained by the hydrometer. In fact, is instrument and a thermometer are indispensable to enable a manufacirer to conduct well any fermenting process. The hydrometer shows that minution of density which measures the gain in alcohol. This attenuation is estimated roughly by the change in the froth or head, which, the presence of saccharine matter is abundant so as to envelope the bonic acid, rises high, but gradually falls as the solution becomes the until, in consequence of the formation of the yeast, a new head rate, for that viscid matter.

## Of the Acetous Fermentation, a Process of Acetification

5957. To acidify, signifies to produce any species of acidity:

tic acid is produced, of which there are several.

5958. Of the processes alluded to, that by which fermented and spin liquids are made to generate acetic acid in the form of vinegar, has designated as the acetous fermentation being accompanied by an apprintestinal reaction between the ingredients in the liquid mixture or so which undergoes this acetifying process. This fermentation differs the vinous in requiring an extraneous supply of atmospheric oxygowhich, as has been mentioned, ethyl is changed into acetyl by the oxid of two atoms of hydrogen, and the acetyl is afterwards acadified by acquisition of two more such atoms (5197), so that, from a hydrated oxide of ethyl, a hydrated trioxide of acetyl arises.

5959. Yet alcohol, whether strong or dilute, does not, per so the change just described. The presence of some substance with attract oxygen from the air, appears necessary to cause its action. Thus dilute alcohol and water do not ferment; but a mixture of the of honey and one of crude tartar to thirteen of alcohol and one heads water, will, in warm weather, produce vinegar in a few weeks (if The change effected in the alcohol may be understood from the fortest of the change 
already given.

5960. The usual method of producing vinegar by the exposure of an open vessels, demonstrates that the necessity of atmospheric organ been learned in practice. Latterly, the process has been greatly expensed by allowing the liquor to fall, in drops, upon the shavings of beach we the temperature being kept up nearly to 100%. According to Lieby, in way one part of spirit of wine, containing eighty per cent, of alcoholy about five parts of water and \(\frac{1}{1000}\)th of yeast of hones or other fermally be converted into vinegar in from twenty-four to thirty-six hours.

# Of the Lactic or Viscous Fermentation.

5961. This has only of late been treated as a distinct process, although effects have long been known to those engaged in the manufacture sugar and fermented liquors, whether for distillation or drink. The manufacture needs in beer, ale, or porter, the premature acidity of the distiller's manufacture referrible to the process under consideration. It is this ferments which supervenes in the absence of yeast, or whenever any nitrogent substance, oxidized by the air to a certain extent, is present. It different the vinous, in giving rise to factic acid, mannite, and a vise matter, usually called ropy, with bydrogen, as well as carbonic a Many years since I was surprised to find the gas given out by eider matter of intense fermentation, take fire, and discovered, on examination, inflammable gas to be hydrogen.

5962. Agreeably to a statement given in Graham, 803, an atom of the

ze, and an atom of lactic acid, are equal to one atom of grape sugar,

nus an atom of oxygen.

5963. I am under the impression that all the four fermentations may sue either successively, or, to a certain degree, simultaneously. Thus, her starch or lactin may be converted into grape sugar. This product ay be partially changed into alcohol, and in part into lactic acid and annite (4074); while a portion of alcohol simultaneously generated, may

undergoing acetification.

5964. Each fermentation has its appropriate ferment. Thus diastase cites the saccharine fermentation, yeast the alcoholic, oxidized diastase, seine or curd, the lactic; while the scum or sediment, called mother of negar, promotes the acetic fermentation. It is the object of the vintner, a brewer, and distiller, to permit only the two first fermentations, the almolic especially, to which the saccharine is accessary. This object is cured by taking great care to have the juice or wort simultaneously substand to a temperature between 60° and 70°, and a limited exposure to air, ith the addition of the proper ferment, where this is necessary; while, by the addition of the presence of any matter capable of inducing the acetus or lactic fermentation is avoided. Much liquor is spoiled by the substitution of the viacous for the alcoholic fermentation.

5965. In a memoir published in the Annales de Chymie, 3d series, 2d ol. 257, Messrs. Boutron and Fremy have made some interesting obsertions respecting the generation of lactic acid in milk. Oxidized caseine 123) is considered by them as pre-emment in efficacy as a ferment, for the lactic fermentation, by acting on the sugar of milk or lactin; but in posequence of an affinity for the generated acid, the oxidized caseine forms

ith it an inert compound which precipitates.

5966. The generation of lactic acid requires the presence both of lactin ad free oxidized casein. Of course, in order to increase the production of se acid, it was found necessary to add an additional quantity of lactin to ilk, but to renew the efficiency of caseine, it was found sufficient to saturate the lactic acid as often as the production of this acid was arrested by

e precipitation of the oxidized casein.

5967. Diastase, after being exposed a few days to the air, becomes caable of inducing the viscous or lactic fermentation. The membranes of se stomach of a dog or calf, or the substance of a bladder, by a like exneure, were found capable of inciting the fermentation in question. Yet aimal matters, in appearance similarly prepared, are productive of different mults, as respects the proportions of mannite, of viscous matter, of lactic zid, or alcohol, generated. The means by which the various ferments, sepectively, produce their appropriate changes are involved in the greatest mourity. Some important additions have been made to our knowledge, as spects the facts. The ferments have all been shown to be vegeto-animal natter in a state of oxidizement, and an analogy seems to have been estalished between their influence and that of some other agents, which have sen considered as acting by what has been called catalysis, which is a ew name given by Berzelius to an old mystery. It has long been known nat there are two modes by which chemical changes are to be excited. no of these, the presentation of one or two extraneous elements causes deemposition and recomposition, by the reactions between the elements so resented, and those subjected to alteration, as in the various cases of elecwe affinity (508, &c.). In the other mode, substances undergo transfornations by being made to rearrange their constituents into one or more new



combinations, by the presence of other bodies with which they do not bine, and which, in some cases, undergo no change themselves it the last mentioned mode of reaction that the name above mentions been applied. Yet, under this head, processes have been crudely and which have discordant features. Liebig indiscriminately gives a complanation to these processes, and to those of fermentation, so for a might be crudely referrible to catalysis.

5968. The following processes are associated by this distinguished mist under one rationale:—the solubility acquired by platina by layer loyed with silver: the catalyzing influence of platina sponge of black: the explosion of fulminating powders by alight causes: the procal decomposition of bioxide of hydrogen and oxide of short agency of nitric oxide in the generation of sulphuric acid: the same

ferments.

tics of the processes thus alluded to. In the case of the plating allow is at least an atom of silver for each atom of plating in actual consists with this metal; and the change which the latter undergoes is process.

same as that to which the former is subjected.

of platina black, causing the acetification of alcoholic vapour, the agent undergoes no change itself; and it enters not into chemical nation either with the materials, or the products. Liebig ascales at the instance to the alternate absorption and subsequent excess oxygen by the powder; since, after exposure to the gas, it may be haustion, be made to give up a portion. But the agency of the mass cannot differ, in this case, from that in which it causes the pure ments of water to combine, and in which, if absorption take place, it is confined to oxygen more than to hydrogen. But the fact established are plate of platina, seems irreconcilable with the idea that absorption are mean of its accomplishment. But if absorption be not operation case, how can it operate in the other?

5971. In this, as in all other cases, Liebig seems to overlank the l important agency of electricity in the phenomena of nature a short infer, that the metal most probably acts by altering the cortect P larity, and consequent association of imponderable matter. But here, assumed, that during the dehydrogenation of alcohol by atmosphere of gen in the presence of platina black, this powder is alternated entered with the power to take it from the air, and to impart it to that of the attraction for oxygen, under the circumstances, is too feeble to air it from the same source, this distinguished philosopher proceeds to make the inference that honey, mother of vinegar, and other substances no motive of acetification, act in the same way by absorbing oxygen for the air, and abandoning it to hydrogen. But if agreeably to the view hore presented, platina black does not act by absorption, no argument, founded on the agency of that substance, will justify the idea that absorption avails if other cases; and it should be recollected, that platina black is very active when perfectly free from moisture, while honey, yeast, mother of views and other substances which cause acctification, have no attraction for an gen in the absence of water: moreover, that the necessity for moisture the preparatory oxidizement of gluten, caseine, diastase, and other orgasubstances, which by exposure in a humid state acquire their capacity

ct as ferments, is inexplicable. Water is powerful both as a catalyzer and a solvent.

5972. Before referring to the absorption of oxygen by honey, as a ground explanation founded on the analogy of platina black, the ability of water

cause honey to absorb oxygen should be first elucidated.

5973. An electric spark or any ignited body, a wire made incandescent y a galvanic discharge, has an influence analogous to platina sponge, of rhich the minutest particle is sufficient to cause ignition throughout an insummable mixture, however large. There is, in this respect, an analogy etween the explosion of inflammable gaseous mixtures and those of gunsween, and of other fulminating powders, of which some, as it is well mown, detonate by percussion or friction, or any cause adequate to derange reduced is the same, whatever may be the exciting cause, and the missest portion of the congeries being made to undergo the change, is of melf competent to produce a like result as respects the whole.

bioxide of lead, have, of undergoing an explosive deoxidizement in conequence of mere superficial contact, is evidently another case, since the rection is reciprocal. In the solution of the alloy of platina with silver, one ody induces another to undergo the oxidizement to which it is itself subseted. In the case of the bioxide of hydrogen and oxide of silver, two odies, both prone to deoxidizement, reciprocally induce that species of bange. But in this phenomena there is no third body to perform a part

malogous to that of the nitric acid.

5975. In case of ferments there is not only the power to produce a hange, but also to produce the particular changes by which sugar, alcohol, nd acetic or lactic acid, and mannite, are respectively generated. Morever, these bodies are themselves undergoing an oxidation or decomposition which is necessary to their power; but this change is not like that which bey induce. Hence, obviously, they operate differently, either from the latina sponge, or platina black, or from the silver in the alloy formed by t with platina. Liebig conceives, that this increased solubility of platina y union with silver, is at war with electro-chemical principles, agreeably which, any metal in contact with another metal, relatively electropositive, becomes less susceptible of attack. But this is not alleged of wo metals in chemical combination, but of masses in contact, or having metallic conductor extending from one to the other. I am surprised that iebig should find the mystery of catalysis lessened by the solution of the Boy alluded to, when it must be evident that if the oxidation of one atom were a sufficient reason why another atom combined with it should be oxilized, an alloy of gold with silver ought to be soluble. Whereas, it is mown that the common process of parting is founded on the utter insoluility of gold when so alloyed.

5976. Liebig alleges that there can be no doubt that the acidification of alcohol is of the same order as the reaction by which nitric oxide provokes be formation of sulphuric acid in the leaden chamber (1019), in which process the oxygen of the air is transferred to sulphurous acid by the intervention of the bioxide of nitrogen, since, in like manner organic subtances associated with spirit of wine, absorb oxygen, and bring it into a

particular state which renders it liable to be absorbed.

5077. But in the case thus cited, for every equivalent of acid formed, an equivalent of the bioxide combines first with an equivalent of oxygen, and

in the next place with an equivalent of the sulphurous acid, for pound which is decomposed by water into sulphuric acid and the bioxide. There appears to me to be no analogy between this that of the influence of matter existing in no equivalent prowhich cannot be shown to form a definite chemical compound acetyl or hydrogen. It is not represented that, in the vinous any union, either transient or permanent, takes place between of the sugar and those of the ferment: on the contrary it is the oxidation and precipitation of the yeast proceeds, pari paralcoholification.

5976. As to all the processes referred to for illustration, as a of fermentation, which they are alleged to resemble, it appears Liebig and his disciples have been too sanguine of their cap

adequate elucidation.

5979. Respecting changes of the kind above described as or Kane uses the following language:—"The elements of a comp tained together in certain molecular arrangement, because t are there satisfied; but it is natural to suppose that whilst remain the same, their affinities for each other might be just a satisfied by a different molecular arrangement." This langue held more reasonably, were this variation in arrangement acct no concomitant acquisition of chemical properties; but is it r consider the difference between sugar, and the alcohol and ca into which it is resolvable, as arising merely from molecular at Can the active influence of alcohol upon the animal nerves be to the situations respectively occupied by its three ultimate posments, carbon, hydrogen, and oxygen, of which it consists? At the union of oxygen with the ingredients of gluten could, by im consequent mechanical impulses, cause the hydrogen and or atom of water to unite with the elements of sugar, and to separat hol and carbonic acid as above mentioned, how can that movem consequent rearrangement of the ponderable particles, explain sition of new properties, of which the combining atoms, or the previously containing them, were destitute? That the presence duces the fermentation of alcohol, and that diastase determines tion of sugar, is admitted; but I am surprised that any philosoconceive, that without first ascertaining upon what the difference perties of alcohol and sugar is dependent, we can understand be ference is caused. Liebig infers that a body in the act of deconcombination, may communicate a movement to the atoms of at compound, so that gluten in the state of oxidation, in which yeast, induces sugar, C12 H14 O14, existing in the same liquid, to the elements of water, making C12 H12 O12, separating into four of carbonic acid and two of alcohol-

5980. Adopting the same views as Liebig, Dr. Kane alleges slow decomposition of diastase communicates to the molecule thousand times its weight of starch, the degree of motion nec their rearrangement, and the appropriation of the elements of waste for the formation of starch sugar."

5981. It is perfectly evident, that the particles of the catalyzed are in some way so affected by the catalyzing body as to be put of reaction, which had not otherwise ensued; but that this is accorded by imparted motion appears to me to be a surmise destitu-

starch is so very small, as is alleged by Dr. Kane, evidently renders it extremely improbable that it acts by creating any mechanical disturbance. Yet this respectable chemist is so completely carried away by this idea, that he proceeds to make the following remark: "This law, of which the simplest expression is that where two chemical substances are in contact, any motion occurring among the particles of the one may be communicated to the other, is of a more purely mechanical nature than any other principle yet received in chemistry; and when more definitely established by succeeding researches, may be the basis of a dynamic theory in chemistry, as the law of equivalents and multiple combination expresses the statical condition of bodies which unite by chemical force."

5982. I perfectly agree in opinion with the author-of these suggestions, as to the purity of the mechanical attributes of the principle on which they are founded, but cannot on this very account deem them competent to ex-

plain the phenomena on which he conceives them to bear.

5983. As the mechanical influence of the motion of bodies is as the weight multiplied by the velocity, is it conceivable that any movement in the particles of one part, by weight, of diastase, can be productive of ana-

logous movements in two thousand parts of starch?

5984. The idea that yeast might owe its power to animalcules, suggested itself to me more than thirty years ago, and seems to have some support from the fact, that fermentation only thrives within the range of temperature compatible with animal life. Latterly, its activity has been ascribed to the power of extremely minute vegetables. Kane, while admitting the existence in yeast of a vast number of globular bodies, possibly animalcules, treats the idea as untenable, because the weight of the alcohol and carbonic acid is greater than that of the sugar employed. But if the union of water with the elements of the sugar, can add to the weight of the products, without the assistance of animalcules, wherefore should their agency be inconsistent with an augmentation from the same source? But the weight of the alcohol and carbonic acid are just equal to that of the sugar, if this be assumed to be in the state of sugar of grapes (5578).

more probable in the case of some species of infection, than in that of fermentation, I conceive that the present state of our knowledge does not allow of our comprehending the means by which bodies, whether organic or inorganic, are endowed with the powers ascribed to catalysis; but that we have great reason to believe that these powers, as well as all the properties which witimate elements acquire by diversity of association, as in compound radicals, are due to the same source as the phenomena of galvanic and statical

electricity.

5986. It is well known, that although pure zinc is not susceptible of oxidation by exposure to dilute sulphuric acid, yet that, when containing misute proportions of other metals, as in the case of commercial zinc, it becomes liable to rapid oxidation by the same reagent. This Faraday explained by the electro-chemical influence of the comparatively electro-negative metallic particles distributed throughout the mass of the zinc, which he conceived to be productive of as many local galvanic circuits with corresponding currents. This explanation has, I believe, been universally sanctioned, and was consistent with the previous discovery of Sturgeon, that when, by amalgamating the surface with mercury, a metallic communica-

tion was made between the electro-positive and electro-negative actility particles, so as to prevent the formation of electrolytic currents through the oxidizing liquid, the zinc became nearly as insusceptible of union with ayen, as when in a pure state.

5987. Nevertheless, either when pure, or when amaigamated, the was found oxidizable by diluted sulphuric acid, provided it were make

element of a galvanic pair.

of the scientific reader, I beg leave to inquire whether the influence the cribed by Faraday to the electro-negative metallic particles has not a great analogy with that of a ferment, than those which have been brought as ward by Liebig, Kane, and others, with a view to explain the influence that class of agents upon mechanical and chemical principles? Wherein may not the distribution of nitrogenated substances throughout a mand inorganic matter, operate as do the metallic impurities in commercial and other trical fishes, shows that the substances which enter into the component animal matter are, when duly associated, as capable as metals of formal the elements not only of simple, but of complex galvanic circuits.

## OF THE PUTREFACTIVE FERMENTATION.

trefactive, animal substances, in general, are much more disposed that petable; and the effluvia which they emit, during the change, are more offensive. It seems as if certain affinities which exist between the ultimate elements of many vegetable and animal substances, although pended by the inexplicable powers of vitality, resume their operation as some as those powers cease, with greater or less activity, according to the more of the substance, and the influence of heat and moisture.

5990. The presence of phosphorus and sulphur contributes greatly to fetor of animal putrefaction. On the other hand, few animal substances of the other hand, fe

susceptible of the vinous or acetous fermentation.

drawn between fermentation proper, and patrefaction. He alleges, that practice the principal mean of discrimination has been the discrimination has been the discrimination has been the discrimination, as attended to the formations, resulting from internal reaction, which are attended to the pleasant smell; whereas fetid processes, in other respects and the property been designated as putrefactive; but that, in point of fact, the present introgen seems to have been the usual associate of substances property is called putrefaction.

5992. But so far as fetidity is an essential attribute of patrefact with presence of hydrogen, with sulphur and phosphorus, seems to me mean sential than that of introgen, since this element is much more rank vehicle of fetid emanations, and, when isolated, is remarkably more rank.

5993. The presence of water, or of its elements, seems indispensive the spontaneous decomposition of organic substances. In no instance either the vinous, aceteus, or patrefactive fermentation and reed stances which are perfectly dry. The effect of desiceat on a present meat and fruits, sufficiently proves the correctness of this allegation. Sprobably, by paralyzing the activity of the water in meat, that sat is a significant probably.

its preservation; and the beneficial influence of sugar upon preserves may in like manner be explained.

5994. The peculiar efficacy of water in promoting fermentation, of whatever kind it may be, rests, as I conceive, on the same basis as its peculiar efficiency in promoting electrolysis. And until we are capable of comprehending the part it performs in the one case, we shall vainly endeavour to

understand the duty which it fulfils in the other.

5995. When, in addition to water, nitrogen is a constituent, the tendency to decomposition is increased. Gluten and yeast, which contain nitrogen, are very liable to an extremely offensive putrefaction. To their deficiency in this principle, Dr. Turner ascribes the indisposition of oils to putrescency; but I conceive their freedom from water, and incapacity to unite with it, to

be the true cause.

5996. The insusceptibility of the vegetable alkalies to decomposition, while containing both hydrogen, oxygen, and nitrogen, may arise partly from their sparing solubility in water, and partly from the predominance of carbon in their composition (5506).

5997. Although heat, to a certain extent, is necessary to putrefaction, it may be arrested by a high temperature, as well as by frost. In the one case, water, being vaporized, is removed; in the other, being congealed, be-

comes inert.

5998. Thenard alleges that water is not decomposed during putrefaction,

but, on the contrary, generated.

5999. Besides water, we may enumerate ammonia, with carbonic, acetic, and sulphydric acid, also carburetted and in some cases phosphuretted hydrogen, among the products of putrefaction.



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### **EMENDATIONS**

ecting the Isomeric Acids of Phosphorus, the Atomic Weight of Silicon, and Composition of Silica.

will be perceived by the readers of that portion of this Comium which treats of "acids relatively to the proportion of base red for saturation" (5181), that a new doctrine has been aded on that subject. Consistently a very important modification een made with respect to the three previously supposed isomeates of phosphoric acid (1153). They are inferred to differ each other only in the proportions of water, or other base h they require severally for their muration; so that there is a basic, a bibasic, and tribasic phosphoric acid (5184). When in ate heretofore designated as free, they are considered as coning three phosphates of water. This assumed constitution of isomeric acids has been represented by Dr. Kane, and other ctable chemists, as affording strong evidence of the existence of ound radicals in certain salfs. Hence having, in arguing against xistence of such radicals, adverted to the constitution of the dift phosphates of water,\* I deem it expedient to give, in the lane of Dr. Kane,† an account of the acids of phosphorus to which ence is made, and of their habitudes with basic water and other

Phosphoric acid has a great affinity for water, combining with it almost exply. It may form three distinct compounds, phosphates of water, the constitution is as follows:—

Monobasic phosphate of water,

Bibasic phosphate of water,

Tribasic phosphate of water,

- PO<sup>5</sup> + HO.

PO<sup>5</sup> + 2HO.

PO<sup>5</sup> + 3HO.

relation was first established by the researches of Graham. Phosphoric acid ies not only with water in these three proportions, but each of them is a type ries of salts, which the phosphoric acid is capable of forming. Thus, there is of monobasic phosphates, another class of bibasic phosphates, and a third, which nost common, of tribasic phosphates; the water contained in the phosphates of being replaced to a greater or less extent, by means of equivalent proportions nonia or metallic oxides.

of water that have been described, and when neutralized by bases may hence totally different salts. The properties of a solution of phosphoric acid may, ore, be totally different according to the manner in which it had been prepared, ence this acid was at one time ranked as a remarkable instance of isomerism; raham has beautifully shown, that the difference of properties is only the result existence of the different states of combination in which the phosphoric acid by exists. It will consequently be necessary to study separately the properties three compounds of phosphoric acid with water.

obasic Phosphate of Water.—A solution of this body reacts powerfully acid, it itates albumen (white of egg) in white curds; when neutralized by a base, it salts which contain but one atom of base, their formula being PO<sup>5</sup> + RO; and ble salt of it produces in solutions of silver, a white, soft, precipitate, PO<sup>5</sup> +

e effort to refute the arguments in favour of the existence in amphide salts impound radical like cyanogen. ements, page 485.

This is the least stable of the phosphates of water, it gradually passes into

the other forms, particularly when its solution is boiled.

Bibasic Phosphate of Water — This form of the acid may be prepared by decomsing bibasic phosphate of lead by sulphuretted hydrogen. It is characterized to combining always with two equivalents of base, forming salts, whose formula is 10 +2RO, its salts give, with nitrate of silver, a white precipitate, PO + 2AgO, which is not pasty like the monobasic phosphate. The salts of this acid may contain only one equivalent of fixed base, the other being water, and may hence, at fest sight, appear to be constituted like the monobasic salts; the basic water is, however, easily known to be present, by its not being expelled by a moderate best with the water of crystallization, but requiring a temperature approaching to ignibe for

its expulsion.

Tribasic Phosphate of Water.—This is the form of phosphoric acid which repres the class of salts most generally known; it is characterized by not precipitates bumon, and by combining with three equivalents of base when fully neutralized. In the majority of cases of the three equivalents of base, one is water; thus the emmon phosphate of soda is a tribanic phosphate, its formula being (PO\* + 2100 BO) + 24Aq; when moderately heated, or even by long exposure to dry air, it less the 24Aq, but it requires to be melted at a red heat, in order to drive off the twenty-file. atom of water, and if this be done, on redissolving the fused mass in water, it eye tallines in a totally different force, and is found to have been changed into blank phosphate of soda, the formula of which is (PO's + 2NaO) + 10Aq. The different is remarkably shown by the action of these salts on a solution of silver; common phosphate of soda precipitates natrate of silver of a canary yellow, and the solut becomes acid; one equivalent of tribasic phosphate of soda, decomposing three equivalents of nitrate of silver, producing one equivalent of tribasic phosphate of silver. two of nitrate of sods, and one of nitrate of water; this last being liquid nitric as The reaction may be simply expressed of course renders the liquor acid

# PO\* + 2NaO.HO and 3 (NO\* + AgO) give PO\* + 3AgO . . . 2(NO\* + NaO) and NO\* + HO.

If on the other hand, bibasic phosphate of soda be used, the liquor remains need for PO's + 2NaO and 2(NO's + AgO) give PO's + 2AgO and 2(NO's + NaO). In the tribasic phosphates, it frequently occurs, that there shall be but one of

valent of fixed base, the other two being water; such salts have frequently as and reaction, and were formerly called hiphosphates. Thus one tribasic phosphate of section PO's + NaO.2HO, the hiphosphate of ammonia is tribasic, its formula being PO's + NH4O. 2HO.

These salts of phosphoric acid were originally designated by Graham, metaphor-

phates, pyrophosphates, and common phosphates.'

It may be proper to add that the opinion of Professor Rose respecting the identity in composition of the different kinds of phosphuretted hydrogen(1166), of which one only is spontaneously itflammable, has been confirmed. According to analysis, either cossists of an atom, or volume, of phosphorus and three atoms, or 43 volumes, of hydrogen, the whole aggregate being condensed atfour.

Their unlikeness, as respects spontaneous inflammability, is ascribed to the presence of impurities which tend either to promote or to retard reaction with atmospheric oxygen.

### Atomic Weight of Silicon and Composition of its Oxide.

In this Compendium (1361), the equivalent of silicon is stated to be 8, and that taking one atom of oxygen to form silicic acid. the equivalent of this compound, known also as silex or silica, is 16 latterly it has been inferred that the equivalent of silicon as 22.22 And that to form silicic acid at takes 3 atoms of oxygen =

Consistently this equivalent of silicic acid is

40 ...

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# BRIEF EXPOSITION

OF

### THE SCIENCE

07

# MECHANICAL ELECTRICITY,

OR

# ELECTRICITY PROPER;

SUBSIDIARY TO THE

### COURSE OF CHEMICAL INSTRUCTION

IN THE

### UNIVERSITY OF PENNSYLVANIA:

WITH ENGRAVINGS AND DESCRIPTIONS OF THE APPARATUS EMPLOYED.

BY

ROBERT HARE, M.D.

PROFESSOR OF CHEMISTRY.

#### PHILADELPHIA:

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John C. Clark, Printer.

1840.



### PREFACE.

Ar the period when the medical school, in which I have the honour to old the chemical professorship, was founded, the science of chemistry exnded to little more than a knowledge of some acids, alkalies, earths, and Its, of which the most important ingredients were unknown. mposition of atmospheric air, and of water, was not understood; the istence of the all-important elements, oxygen, hydrogen, and nitrogen, t having been detected. The subsequent discovery by Dr. Black, of the tent agency of heat, was the first step in a department of our science, hich at this period occupies, in a course of chemical lectures, a large poron of the time. The discovery of the same great chemist that atmospheric r was not to be considered as a fluid sui generis—that there might be more an one kind of aëriform matter; by leading to a knowledge of the gases, lded another topic, which, in all its bearings, practical and theoretical, may considered as no less entitled to attention, than the phenomena of heat. The invention of the Voltaic pile, and the employment of its wonderful wers in effecting chemical decomposition, besides leading to a knowledge many chemical facts and agents, connected either in theory or practice, e whole science of electricity, whether galvanic or mechanical, with cheistry.

To these topics have latterly been added the electromagnetic properties of atter; which by their association with electricity, with a power of chemilar decomposition, and of giving shocks to the animal frame, force them-lves upon the attention of the chemist and physiologist.

Both the theory and nomenclature of our science, are at present grounded son the electrical habitudes of chemical agents. Under these circumances, a knowledge of galvanic and mechanical electricity is of fundaental importance; and to impart this knowledge, where it has not to a efficient extent been made part of the student's previous education, should a primary object.

These impressions are sanctioned by the practice of the most distinuished writers of elementary works upon chemistry, in each of which I elieve a brief treatise, on the electrical sciences, is made to precede that pon chemistry proper.

#### PREFACE.

on in lachool, by the fact, that it if as i it remedies. The question who or to will a be presented to the practition, to it of their nature, or the mean ir influence on the vital function of y.

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# A BRIEF ACCOUNT

If the Origin of the Science of Electricity Proper; and of the Progress which it made prior to the time of Galvani.

1. An attractive power is acquired by resins, sulphur, glass, and a variety of other substances, when rubbed; and if the masses, thus excited, be sufficiently large, the phenomena of light, of mechanical concussion, and ignition, may result, and even a feeble imitation of thunder and lightning.

2. As it was in amber, in Greek called *Electron*, that the ittractive power, arising from friction, was first observed, he principle to which it was ascribed was called electricity, and all substances, in which it could be produced,

electrics.

- 3. We are informed that Thales of Miletus, who flourished six hundred years before Christ, was so much struck with his effect of the friction of amber, as to imagine that it night be endowed with animation. Subsequently, it was secretained that the attractive power, which had been observed in amber, after being rubbed, might be produced, by he same means, in other resinous substances, and in the ourmaline, or lyncurium, as it was then called, by exposure to heat.
- 4. No further progress was effected in electrical knowedge until the seventeenth century, when, after considerable additions had been made to the catalogue of electrics by Gilbert and Boyle, Otho Guericke discovered that light and sound might result from electric excitement. His observations were made by means of a globe of sulphur, cast n a glass vessel which was fractured to extricate the casting. Little was it suspected by the ingenious operator, that the glass globe, thus broken, would have answered better for the purpose in view, than the globe of sulphur in the moulding of which it was sacrificed.
- 5. The discovery of the usefulness of glass as a mean of producing electricity, appears to have been made by

Hawkesbee, we owe the remark that the electrical excuement of glass and other electrics was communicable to other bodies when insulated, not only by direct contact but by wires or threads of great length; and by the electrician, in conjunction with another named Wheeler, it was first observed that this property of conducting the electric virtue, while belonging to flax or hemp, did not belong to silk; also, that, by the class of bodies in what electricity can be excited, it cannot be conducted; while in those by which it may be conducted, it cannot be creited. Thus were two classes of bodies distinguished: one as electrics, or non-conductors; the other as non-cleotrics, or conductors.

6. It was ascertained, however, that a conductor, if supported by a non-conductor, might receive the electric true from an excited electric. A conductor, so supported

was said to be insulated.

7. Du Faye soon after ascertained the important trub that there are two kinds of electrical excitement. One of these, being observed in glass, was called vitreous; and the other, resinous, because observed in resins. By the communication of either species of excitement, light bodies excited by means of resins were attracted by such as were excited by means of glass; and when these opposite excitements were made, in due proportion in different insulated conductors, on bringing the conductors together. I neutralization, and of course an apparent annihilation, of the electricity in both was the consequence.

8. The means of collecting the electric fluid were some after much improved in Germany; where sparks sufficient to kill birds, and ignite spirits and other inflammable matter were produced by the joint influence of several globes.

simultaneously excited.

9. In the year 1746, the Leyden phial was invented. Cuneus and Mushenbræck, attempting to charge the water contained in a phial with electricity, a shock was experienced by Cuneus, who happened to touch the conductor with one hand, while grasping the phial with the other.

10. This phenomenon was soon after explained by Franklin. He had ascertained that, whenever either kind of electricity is communicated by friction to one surface, the other kind will be created in that by means of which the friction

is effected, provided it be insulated.

11. When a glass is rubbed by the hand, it takes electricity from the hand, and from the person to whom it belongs. If standing on a non-conductor, the person will be electricited, at the same time that the glass which he rubs may, by due proximity, excite another body; but, then, the electricity of the person who rubs the tube, and that of any body to which the tube may electrify are of opposite kinds; the former being the resinous, the latter the vitreous electricity of Du Faye. A stick of resin would cause the opposite result, producing vitreous electricity in the person rubbing it, and resinous in a body touching the resin, subsequently to its exposure to friction.

- 12. These phenomena were thus accounted for by Frank-Some bodies by friction (glass for instance) acquire additional power to hold the electric fluid, and hence draw it from the conducting body rubbing them, and give up the excess to any adjoining conductor, when the friction Resins, on the other hand, have their capacity for the electric fluid lessened by rubbing, and hence, while subjected to this process, give it out to the rubber, and afterwards draw on any adjoining body to supply their deficiency. Glass and resin, therefore, produce both kinds of electricity, which are merely the result of an accumulation, or deficiency, in an insulated body, of a fluid which pervades the creation. A conductor, charged in either way, will produce an electrical current when presented to other bodies in connexion with the earth. In the one case, electricity will flow into the conductor; in the other case, it will flow out of it.
- 13. Franklin, also, discovered that, when an electrical stream is directed into a phial, situated like that of Cuneus, there is, at the same time, a stream proceeding from the outside; so that, in proportion as one surface gains, the other loses; and, accordingly, in a charged phial, one surface will be found vitreously, or redundantly, excited, the other, resinously, or deficiently; and a light body, after touching either surface, will be repelled by it, and attracted by the other.
- 14. He inferred that there was only one electric fluid, to different states of which, the names of vitreous and resi-

nous electricity, had been erroneously applied. The latter

he called negative, the former positive electricity.

15. Franklin afterwards identified lightning with electricity, by drawing this fluid from the clouds by means of a kite; availing himself of a contrivance, which had previously been appropriated to juvenile recreation, to make a most sublime and useful discovery.

# MECHANICAL ELECTRICITY,

OR

# **ELECTRICITY PROPER.\***

#### ORDER TO BE PURSUED IN TREATING OF ELECTRICITY.

DESCRIPTION OF ELECTRICAL MACHINES.—Usual means of producing Electricity.—Communication of Electricity.—Different kinds of Electricity.—Means of accumulating Electricity.—Means of accumulating Electricity.—Means of Electricity.—Effects of Electricity.—Additional Means of producing Electricity.—Theoretic explanation of Electrical Phenomena.—Means of electricying Patients either by sparks or by shocks.

### GENERIC DESCRIPTION OF THE ELECTRICAL MACHINE.

- 16. I am unacquainted with any apparatus, designated as an electrical machine, which does not consist of an electric, so situated as to be conveniently subjected to a friction calculated to produce electric excitement, one or more collectors, attached to a prime conductor properly insulated, and one or more cushions, for rubbing the electric. The cushions, in the more perfect forms of the machine, are associated with another insulated conductor.
- 17. Experience has shown that, of all the electrics, glass is the best for the construction of electrical machines; and as of all the possible forms, only two are much in use,
- "It seems to be impossible to designate well by any appellations heretofore employed, the different branches of electrical science. As almost all the phenomena ascribed to Electricity, before the time of Galvani, are due to mechanical causes, I have heretofore concurred with some electricians, in distinguishing the branch under which these phenomena are treated, as "Mechanical Electricity;" but it may be objected that the discovery of the magneto-electric machine, and other magneto-electric contrivances, causes another set of phenomena to be produced by mechanical causes. This objection may be mitigated by the fact, that the phenomena created in the last mentioned way, are not the consequence of mechanical causes simply, but of the reiteration by them, of the influence of a magnet. I have, however, subjoined another appellation, "Electricity proper," which I should substitute for that first mentioned, were not the retention of a defective name often less disadvantageous than employing a new one, however unexceptionable.

those of the cylinder and of the circular plate, it may be inferred that these are preferable.\* In either case, the fretion is produced by the rotation of a shaft, occupying the axis of the cylinder or plate, and fastened by screws a cement. The shaft or axis, being secured in one or more collars like the mandril of a lathe, is turned by a wind. or by a band and wheels, which are so proportioned and arranged as to quicken the motion. In order to prevet that access of the air to the surface of the glass, which causes a loss of electricity, it is usual, besides the cushon to have a silken flap, of which one border is sewed to the edge of the cushion, so that it may extend from the cushion, till the other border approximates the collectors

#### Electrical Machine, with a Plate four feet in diameter.

18. The opposite engraving represents a machine, constructed so as to be purposently affixed to the canopy over the hearth of my lecture room.

19. The convenience of having this machine in the situation which it secure.

have found to exceed my most sanguine expectations; as it is always at hard je never in the way In lecturing, with the aid of a machine on the mine level of the lecturer, one of two inconveniences is inevitable. Either the machine will casionally be between him and a portion of the audience, or he must be between portion of the audience and the machine. Situated like that which I am about b describe, a machine can neither hide the lecturer, nor be hidden by him. With it its power at his command while kept in motion by an assistant, he has no part of a to reach or to handle besides the knob and sliding rod of the conductor, which are the most convenient situation.

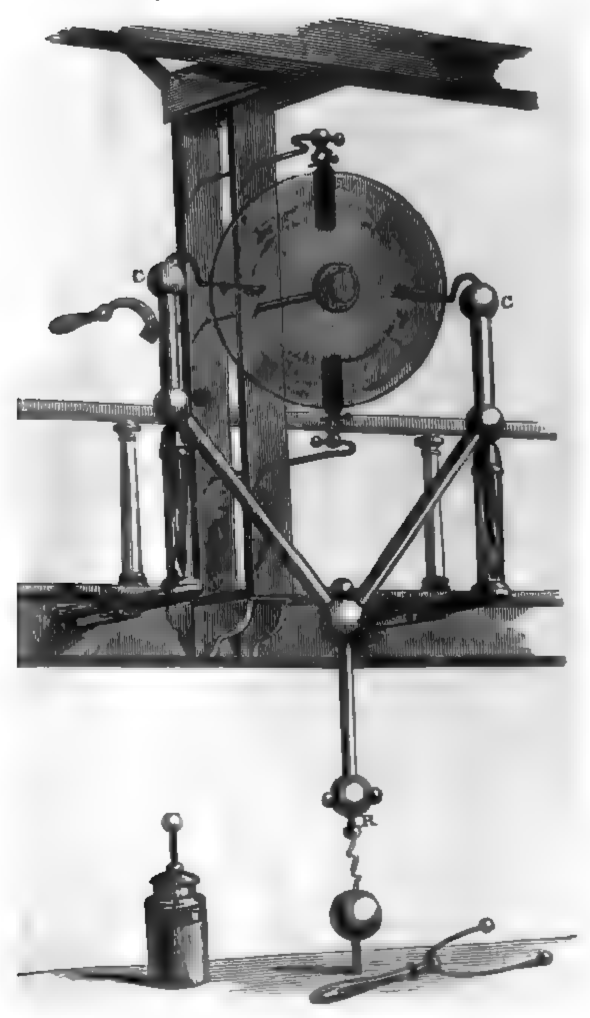
20. The object of this machine being to obtain a copious supply of electronic to experiments in which the positive excitement is always employed, it was not does necessary to insulate the cushions and the axis

21 The prime conductor is insulated by the same means as those of the borne tal plate machine. (c. 27) At C C are the collectors. R represents a chine tal, which may be drawn out to such an extent, as to be brought in contact which apparatus placed under it upon the table. The cushions are, in their constraint, analogous to those described as belonging to the horizontal plate machine, and to presented by Fig 3, (c 29.) Their situation and mode of their application, most to evident from the representation of them in the figure.

#### Engraving and Description of an Electrical Cylinder Machine.

22. A, the glass cylinder—C, the positive, or prime conductor, supported to a glass pillar—E, the collector with its points so projecting as to be quite near to the cylinder. The negative conductor, F, is also upheld by a glass pillar, supporting the rubber or cushion to which the silk flap, G, is attached. By means of a set

<sup>\*</sup> I am under the impression that the great disadvantage attendant on the ow of cylinder machines, is the presence of moisture within the cavity, which, being moccasible, cannot be desiccated. This moisture enables the glass to hold the charge as in a Leyden jar, so that it does not relinquish it to the collectors. The evil the arising, may be obviated by taking great care, in the construction of the machine remove all the moisture by a blast of air rendered anhydrous, and forthwith closes the apertures in the necks, by putting on the caps air-tight. But to keep out more ture, the caps must be of metal, since, if constructed of wood, moisture will prestrate. Moreover, the coment after a time is liable to become leaky. I concerve that it is in consequence of the presence of moisture, that some cylinder machines, which otherwise will not give a spark, perform well after being heated by a bot was or a pan of coals. These remarks are verified in the case of the cylinder machine. of which an engraving and description is given in the opposite page.

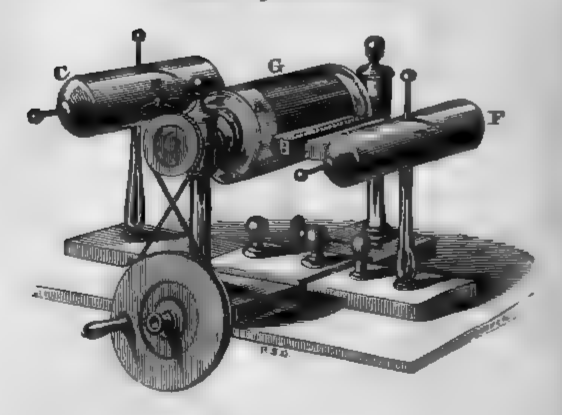


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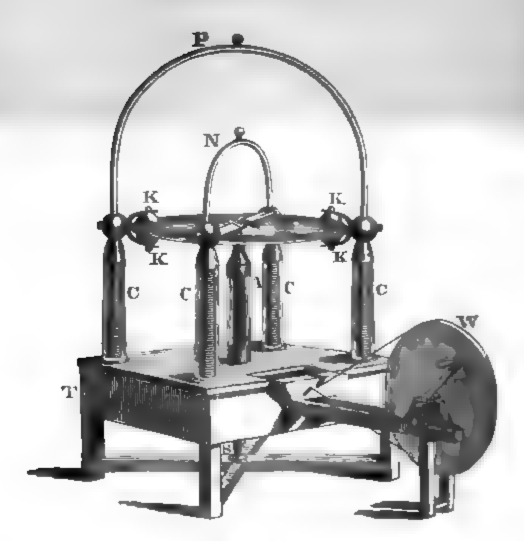




## Electrical Cylinder Machine.



Electrical Plate Machine.



(Page 7)

w, the larger wheel may be made more or less remote from the smaller one, so adjust the tension of the band by which motion is communicated to the one the other.

The former being turned by the winch, causes the smaller one to revolve, and surse the cylinder to the axis of which it is affixed. The revolution of the der while the cushion is pressed against it by a suitable spring, causes the frictwhich is requisite to excite each portion of the cylinder as it successively passes sushion. Each of the conductors is furnished with projecting brass knobs, ace to take sparks.

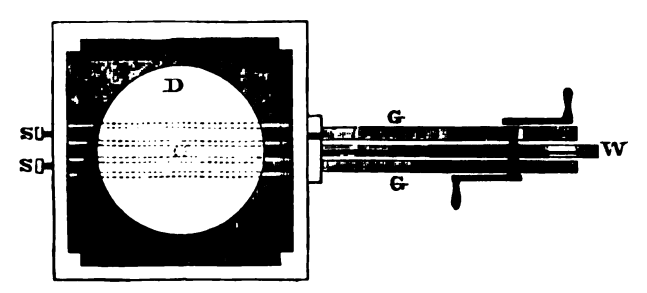
# ription of an Electrical Plate Machine, the Plate mounted horizontally, and so as to show both Negative and Positive Electricity.

The power of electrical plate machines has been generally admitted to be ter than that of machines with cylinders. The objection to the former has been difficulty of insulating the cushions, so as to display the negative electricity. pting the plate machine contrived by Van Marum, I have read of none in h this difficulty has been surmounted. It is still insisted upon by respectable ricians, as if it had not been sufficiently removed by his contrivance.

. I shall now describe a plate machine, by which both electricities may be pro-

d, and which I have used successfully for eighteen years.

. The plate, B, (thirty-five inches in diameter) is supported, as represented in the e, upon an upright iron bar, about an inch in diameter, covered by a very stout s cylinder A, four inches and a half in diameter, and sixteen inches in height, only at the base, through which the bar is introduced, so as to form its axis. summit of the bar is furnished with a block of wood, turned to fit the cavity, ed at the apex of the cylinder, and cemented therein. The external apex of the ider is cemented into a brass cap, which carries the plate. The glass cylinder ble to no strain. It is only pressed where it is interposed between the block of 1 within, and the brass cap without. The remaining portion of the cylinder s only its own weight, while it effectually insulates the plate from the iron axis. brass cap is surmounted by a screw and flange, by means of which, a corresling nut, and disks of mahogany, the plate is fastened. A square table serves basis for the whole. The iron axis, descending through the top of the table, is ished with a wooden wheel of about twenty inches in diameter, D, (Fig. 2,) and inates below this wheel in a brass step, S, supported on the cross of wood, which the legs of the table diagonally together. The wheel, D, is grooved and made to lve by a band, which proceeds from around a vertical wheel, W, outside of the This external wheel has two handles, by means either of one or both of th it may be turned. It is supported on two strips of wood, G G, which, by opriate screws, (represented at S S, Fig. 2.) may be protruded, lengthwise, from s, which confine them from moving in any other direction. Consequently, the nce between the wheels may be varied at pleasure, and the tension of the band sted.



Nearly the same mode of insulation and support, which is used for the plate, is I in the case of the conductors. These consist, severally, of arched tubes of s, of about an inch and a quarter in diameter, which pass over the plate from side of it to the other, so as to be at right angles to, and at a due distance from, other. They are terminated by brass balls and caps, which last are cemented on s cylinders, CCCC, of the same dimensions, nearly, as that which supports the e. The glass cylinders are suspended upon wooden axes, surmounted by plugs ork, turned accurately to fit the space which they occupy. The cylinders are

surrounded and secured below, by wooden rings acrowed to the table. In the conductors are effectually insulated, while the principal strain is been wooden axes.

28. The form of the collectors (c. 16,) must be sufficiently evident from the centation of them in figure 1st. They consist of hollow hemispheres of the within which several points proceed towards the plate from their centres may be when they are attached to the knobe, kkkk. The hemispheres are attached to the knobe, kkkk.

29. The cushions (c. 16.) are represented by Fig. 3. These are included the springs, by which they are made to press with an elastic force upon that of the glass, the degree of the pressure being regulated by the screw, S.

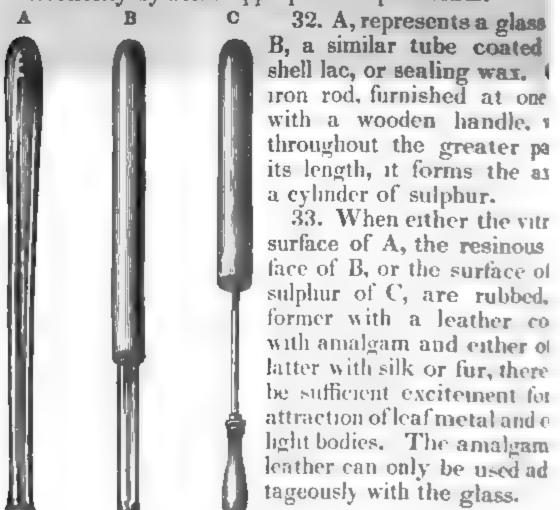


#### USUAL MEANS OF PRODUCING ELECTRICITY

30. It has been stated, in the preceding remarks an attractive power is acquired by resins, sulphur, and a variety of other substances when rubbed:—also bodies, susceptible of this species of excitement, are electrics, and the principle, on which it is supposed dependent, is called electricity.

31. I shall proceed to explain and illustrate the evol

of electricity by some appropriate experiments.



n T

# Experimental Illustrations.

34. Attractive power imparted by friction to amber, plass, resin, sulphur. Large glass tubes rubbed—also, ylinders of sulphur and resin. Thin metallic leaves attracted, at the distance of several inches, either by the plass, the sulphur, or the resin. Electrical machines put n operation.

### COMMUNICATION OF ELECTRICITY.

- 35. The electric virtue cannot pass from one part of an electric which is excited, to another part without extraneous aid; nor can it pass off, from one electric, through any other. Hence these substances are called non-conductors. Through metals, on the other hand, it escapes instantaneously. It passes through flax or hemp, but not through sik. Water it pervades with great facility, or any thing which contains moisture.
- 36. Substances which are thus capable of transmitting electricity are called conductors, and are divided into perfect and imperfect conductors. The metals are the only perfect conductors. All other conductors are imperfect; and, at the head of this class, is charcoal, as being the best conductor of electricity next to the metals.

Experimental Proofs that Electricity cannot pass from one part of an Electric to another, through, or by means of, the Electric.

# An Experimental Illustration.

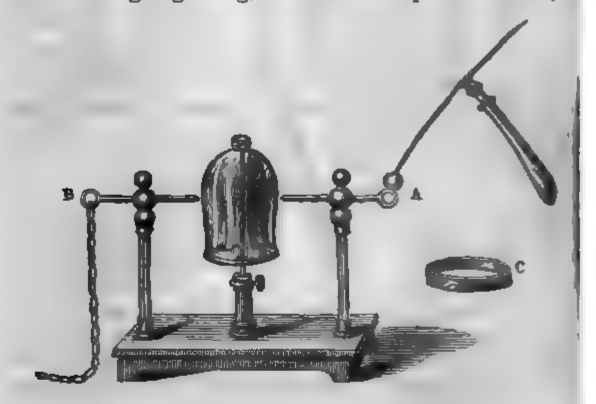
37. One part of a mass of glass, of sulphur, or of resin, being so excited by friction, as to attract light bodies, another, not being rubbed, is not found to attract them; so that the electric virtue does not pass from one part of the mass to another.

# Peale's Experiment.

38. Among a great abundance of ingenious electrical contrivances, that which we owe, as I believe, to Mr. Franklin Peale, deserves more than ordinary praise for its simplicity of construction, and beauty of effect. One

B

One modification of Peale's apparatus is represented the following engraving, which I shall proceed to explain



between wires supported on glass pillars. One wire, A, community with the positive, the other, B, with the negative pole of the made Under these circumstances, if the machine be put in operation, the p tion of the glass next to the wire, A, becomes positively excited, that a B, negatively excited. Consequently, agreeably to the general law t bodies similarly electrified separate, those dissimilarly electrified appropriate each excited portion of the glass will move away from the wire such excited, and will seek that which is differently excited. Thus the remains changing the situations of the excited parts, and their exchange situation reversing their excitements, a rapid movement must ensue, a only mode in which it can take place freely; I mean that of a relative the pivot.

40. The bell is ornamented by strips of gilt paper, which renders motion more sensible to the eye; but no coating is requisite to the app

priate effect. A bare bell glass is sufficient.

41. If the metallic band, C, be made to encompass the bell glass, it is be found incapable of receiving any motion from the electrical excitone. In fact, the electric fluid will be seen passing into it on one side, and paing out of it simultaneously on the other side; proving that it can under these circumstances, retain any excitement, in consequence of the side of the consequence of the c

conducting power of the metallic band.

42. Mr. Peale's experiment was performed by means of a globular givessel with a short neck, or perforation for the admission of the pivot on it will be perceived that I have availed myself of this apparatus, to sho that the electrical excitement communicated to one part of a non-conduct does not extend itself to others; and that consequently in different pursuithe same non-conducting mass, opposite kinds of electrical excitement of the produced. It is owing to this property, that the glass bell is put the motion. The electric fluid, being unable to pass along the glass, in its class.

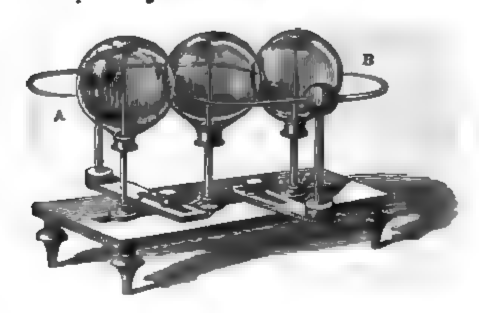
sek an equilibrium, moves the glass along with it, in consequence of an ection arising between each wire, and that part of the glass which has Terent excitement.

### Experimental Illustrations.

13. That the phenomenon owes its existence to the nonducting power of the glass, is shown by encircling the with a metallic band while subjected to the action of machine as above described. Under these circumnces, as the fluid passes from one side of the glass to other with perfect ease, that diversity of excitement, ich would cause the rotatory motion, cannot arise.

#### Modification of Peals's Experiment.

4. The following figure shows a modification of Peale's apparatus, by which I e contrived to put three globes in motion.



i. The arched wires. A, B, being severally connected with the different conductof an electrical machine, cause opposite states of excitement upon the surfaces be globes, in the neighbourhood of their pointed terminations. A revolution of excited surfaces, arising from the attraction of the differently excited wires, consently ensues, in a manner analogous to that described in the case of the bell abovementioned.

#### Additional Illustration.

3. The principle explained by the preceding experiment receives another asons.

ing engraving is intended.

7. A glass ball is supported on a glass plate. On the plate, strips of tin feil are ested, as to form a broad circle or border near the margin of the plate, and four i to that circle. There is likewise a flat brass ring, supported and of course inted by glass pillars, so as to have its inner edge immediately over the exterior a of the feil.

8. The brass ring being in communication with the prime conductor of the maie in operation, and the tin foil in communication with the cushions, the ring foil will be oppositely electrified. The ball, being attracted by the ring, becomes tively electrified in the part which comes in contact with it. The part thus elected will then be attracted by the foil, and communicating its thurge, return to



h

the ring to undergo another change. Different parts in succession undergo the electrical changes, and the consequent movements are, of course, complement 49. Peale's apparatus is, however, preferable to this which has just been a scribed, as it is more easy to succeed in making it perform well.

Experimental Proofs that Metals, Charcoal, Moisture, and Flax or Hemp are Conductors of Electricity, and that Sulphur, Resins, Glass, Silk, and Wool are Nonembuctors.

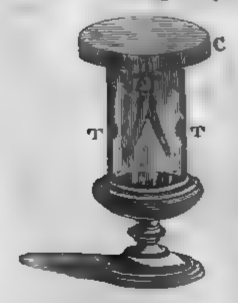
- 50. The electrical machine, being in operation so as to emit sparks, or to act upon pith balls, or other light bedies, those electrical effects cease, when the conductor, which is the immediate cause of them, is touched with a rod of metal, or by a piece of charcoal, communicating with the earth; and they are enfeebled, when one end of a hempen or flaxen string is attached to the conductor, while the other end is held in the hand, or lies upon the floor.
- 51. The conductor being excited, as above described, the phenomena do not cease, in consequence of the contact of a glass rod, or of cylinders of sulphur or resin; nor are

rings, as in the case of those of flax or hemp. The lass rod, or the woollen or silken strings, being moismed, the electricity is carried off by them from the conactor. It escapes also readily through a tube filled with rater.

### DIFFERENT KINDS OF ELECTRICITY.

- 52. It may be learned, from the brief account of the ise and progress of electricity, (e. 7) that the electrical xcitement which may be produced in glass, by friction, liffers from that which may be produced, by the same neans, in resin, or sulphur: that light masses, as paper, or pith balls, separate from each other, when either excitement has been imparted to both: but if one body receives the resinous, the other the vitreous excitement, an attraction between them will ensue. Both excitements, in due proportion, neutralize each other. Also, whenever either excitement is produced, in one body, the other will arise in some other, if both bodies be supported by nonconductors, so as to prevent the escape of electricity as soon as generated. Hence if a person, standing on a glass stool, rub a tube of the same material, he will be found resinously electrified; while any body to which the glass may be presented will be vitreously electrified. A stick of resin, being substituted for the glass, and rubbed in like manner, and under the same circumstances, the same phenomena will appear, in a different order. The person rubbing the resin will be vitreously excited; while the excitement of the body to which it is presented will be resinous.
- 53. In like manner, when the cylinder of the electrical machine is put in motion, the insulated cushion which rubs it, acquires the resinous excitement, while the prime conductor becomes excited vitreously. If a globe of sulphur or resin were substituted, the cushion would receive the vitreous excitement, while the conductor would be excited resinously.

### Description of Bennet's Gold Leaf Electrometer.



54. A glass cylinder, supported by a make tallic podestal, is surmounted by a make canopy; from the centre of which, two and ing strips of gold leaf are suspended. Structure of tin foil, T T, are pasted on the glass, we to terminate at the upper ends a latter the level of the lower ends of the gold and and so as to be in contact below, with a pedestal. This should be uninsulated. The gold leaves are more energetically attraction consequence of the proximity of the terminal foil.

## Experimental Illustrations.

55. The leaves of the electrometer diverge on the proach either of excited sulphur or glass; but when but are approximated to it at once, the leaves will display divergency.

56. Electrical machine shown, each conductor being furnished with a quadrant electrometer. As the plate is turned, the pith ball of each electrometer rises. As often as a spark is taken from either conductor, the pith ball of the electrometer associated with it falls; and when a metallic wire is made to touch both conductors simultaneously, neither of the pith balls indicates any excitement.

# Of Electric Poles and Electric Circuits.

57. There is a resemblance, which will hereafter be recurred to, between the reciprocal action of magnets, and that of electrics of which the extremities are in opposite states. Under such circumstances, the poles similarly electrified, or similarly magnetised, appear to repel each other; while those in dissimilar states appear to exercise a reciprocal attraction.

58. The extremities of the magnets, from their exercising a reaction with the terrestrial poles analogous to that which they exercise reciprocally, were called poles: that which is attracted by the north pole of the earth being

From the analogy, the extremities of excited electrics, and of galvanic and voltaic instruments having properties resembling those of excited electrics, received the same appellation. Hence electricians speak of the positive and sthe negative poles, of the electric machine, of the galvanic statery, or of a voltaic series, whether in the form of the trough, or of the pile.

those parts of an electric, galvanic, or voltaic circuit, at, or between which, ignition, light, chemical decomposition,

Tor sensation are perceived.

5

- communicate with each other through a wire or other competent conducting body or bodies, an electric circuit is said to be formed. The electric fluid flowing into the cushion from the negative conductor supporting it, is by the electric carried to the conductor which supports the collector, whence, by means of the conducting communication, it returns to the negative conductor. Agreeably to the hypothesis of two fluids, there is in such case a double circuit. Two fluids, separated from each other by the friction, move in opposite directions, and meeting and combining in the conductors as soon as separated, the equilibrium is not sensibly altered.
  - 61. In making or breaking such a circuit, sparks will appear at the time and place, when and where the interruption is such as to allow them to occur. In that case the poles are at the points between which the sparks pass, or indications are afforded of electrical agency.
  - 62. The poles of an electric machine and the conductors are often mentioned as if they were identical. In fact the poles are usually at some points in the surfaces of those conductors. They vary, however, at the pleasure of the operator; since they always exist at those parts of the conductors (or of any conducting body or bodies touching one or both) which are nearest to each other, or at which sparks pass.

63. It should be understood that any conductor, especially any perfect conductor, assumes the same electric state as the pole, with which it may be in connexion; and that a contact between either pole, and one extremity of a rod, or wire, transfers the polar influence to the other

at which the excitement is most active, and expenses shows this state to exist always, either at the most proment part of the electrified mass, or that nearest to any

prominent part of the other pole.

64. In operating with an electric machine, of which the rubber is uninsulated, the earth usually forms a per of the circuit, and any conductor, the human body for the stance, which, while it has a conducting communicated with the earth, happens to be nearest to the positive polacts as the negative pole, when the machine is in operation, and there is sufficient proximity. This is indicated by the passage of sparks.

65. I have dwelt upon this subject more particular the acceptation of the word pole among men of science latterly, by its practical and theoretical associations become of the highest importance. (See my Treatise at

Electro-magnetism, page 78.)

66. Since the preceding explanation was published, it raday has suggested that electrical poles as above defined should be called electrodes; the positive pole being called the anode, the negative pole the cathode. (See my Treats on Voltaic Electricity, page 7.)

### MEANS OF ACCUMULATING ELECTRICITY.

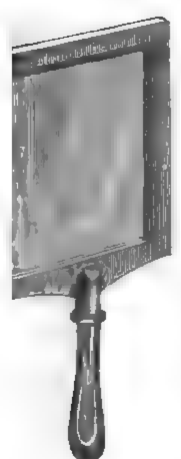
67. In the case of the insulated conductors of an electrical machine, oppositely excited by the revolution of a glass surface which successively rubs the cushions supported by one of the conductors, and passes under metalic points projecting from the other, it has been shown that a is only necessary to make a communication between them by a perfect conductor, in order to destroy their respective excitements. It follows that the surcharge in the one must have been just equivalent to the deficiency in the other; and that of course the whole quantity of electricity in the conductors is the same, whether they be in a state of excitement or of neutrality.

68. It has also been shown, that the electric fluid does not pass through electrics, or from one part of an electric

to another.

69. I shall now proceed to demonstrate that, if an electric, sufficiently thin and strong, as a pane of glass for astance, be charged on either side with either kind of elecity, the other side of the pane will acquire proportion, a charge of the opposite nature.

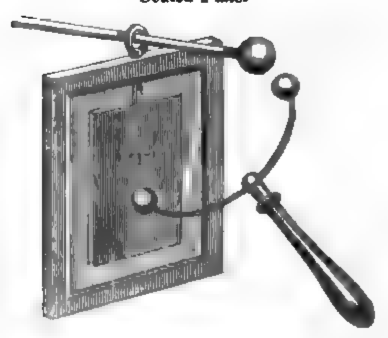
#### Experimental Illustrations.



70. A dry glass pane being held in one hand by its insulating handle, sparks are taken from the excited conductor of an electrical machine, with the knuckle of the other hand. The glass pane, being interposed between the knuckle and the conductor, at first does not appear to intercept the sparks; yet, as they gradually diminish and finally cease, they are obviously intercepted by the pane, sooner or later. The pane being supported by the handle, on touching the surfaces in the part which has been exposed to the sparks with one hand, while the part opposite, on the other side of the pane, is touched by the other hand, an electrical discharge takes place, a shock is experienced, and the electrical excitement disappears.

71. It follows, that the sparks which had apparently passed through the pane, had actually been arrested by the surface nearest to the conductor, and had appeared to reach the hand, because, for every spark received on one side, an equivalent portion of the electric fluid was expelled in the same form from the other side.

#### Coated Pane.



2. A pane of glass having been coated on both sides with tin foil, upting a space of about two inches from the edge all round; the coatare made severally to communicate, for a short time, with the two lated conductors of an electrical machine, while in operation, the coatbeing otherwise insulated. Thus situated the pane becomes charged electricity. The different coatings being, in the next place, allowed communicate with each other through a metallic arc, a discharge en-

sues, more or less powerful, according to the power of the machinextent of the coated surface, and the dryness of the air.





73. In the "Brief Account," in (e. 9, &c.) some account has being this celebrated invention. The cut is a representation of it in an approximation. There is no essential different tween a coated phial, and a contribute existence or absence of curve the interposed stratum of glass, do sensibly affect the result in them in practice excepting as respects in iency.

74. The form of the jar is more! able to the retention of a charge does not allow that free intercess tween the inner surface and the sit, carries off the electricity, pan pass

each coating.

75. The coating, it will be observe represented as extending both on the and on the outside, till within a different the brim equal to about one

of the whole height of the jar. The jar is closed by a broad a stopple, through the centre of which passes a metallic rod, terminated by a knob, and below by a spiral of wire, which establishes a secontact with the inner coating.

76. The most convenient mode of charging such a jar is to grant the hand, and present it to the knob of the prime conductor, who cushions are uninsulated, or in communication with the outer communication of a chain or wire.

### Of the Common Discharger.



- 77. The instrument, represented by the foregoing figure, is called a disharger, being employed to complete the circuit between the oppositely harged surfaces; thus enabling the excess in one to be discharged upon he other.
- 78. The rods, R, R, are joined by a hinge; so that by means of the flass handles to which they are severally affixed, the terminating knobs may be made to come in contact, or to be more or less remote from each ther.
- 79. A charge is imparted equally well by the contact, or communication of either coating with either conductor.
- 80. The charging of a coated pane or jar may be effected, provided either of the surfaces are in communication with either conductor, the other surface communicating with the other conductor, one or both of the conductors being insulated. In whatever way a charge may be induced, it will be found that the one surface loses as much as the other gains; since a conducting communication is always sufficient to bring them both back to a state of neutrality. Hence as the sum of the quantities on both sides of the pane is always the same, charging the pane does not derange the electrical equilibrium of the surrounding medium.
- 81. The charge must vary in its strength according to the power of the machine, the dryness of the air, and the thickness of the glass; since the self-repellent power of the electric particles, and their attraction for the negative surface of the glass, must be inversely as the squares of

the distances at which they operate.

82. On this account, as a plate of mica may, with less thickness, possess greater strength than a glass pane, it will, in proportion to the respective areas, receive a much higher charge. A certain degree of strength is necessary to enable the glass to resist the intense attraction between the surcharge of electricity on the positive side, and the surface of the glass on the negative side. In obtaining this strength in the glass, we increase the distance between the surfaces, and of course diminish the efficacy of the self-repellent and attractive powers on which the charge depends.

## Experimental Illustrations.

83. That the charges may be imparted through either coating by either conductor, shown by duly charging and discharging coated panes and jars.

Experimental Proofs that the different Surfaces of a d Electric are oppositely electrified.



84. If, from the conductor electrical machine in open Leyden jar be suspended by tallic hook, connected with its coatings, it receives no until the other coating is a mated by a conducting sub which communicates directly directly with the other coating of the machine.

85. If one or more bodi for instance, little metallic qualified to act as bell clapp suspended at a suitable d

between bells severally communicating with the coof a charged pane or jar, the balls will play between bells, demonstrating them to be oppositely electric their communication with the different surfaces.

Chime of Bells.



86. The little metallic balls, b, b, and the bell situated between them, being suspended by silk, which is a non-conductor, cannot receive the electrical excitement mmunicated to the inner coating of the jar. But the two outer bells being suspended by metallic chains from a metallic rod communicating with the inner coating, will partake of the excitement in this coating. Consequently the balls remaining neutral, while the outer bells are excited, are attracted by these, and on coming in sentact with them receive a quantity of the electric fluid adequate to bring them to the same degree of excitement. Hence in the next place they separate from these bells, and are attracted by the central bell, which, by means of a chain connecting k with the external surface of the jar, is brought into the same electrical state as his surface, and must of course be in a state opposite to that of the bells communinating with the other surface. The balls, by contact with the central bell, lose as nuch electricity as will bring them to the same state as this bell, and are then separated from this bell; and are attracted by the others, are again separated, and are ugain attracted, until by repetition they transfer the surcharge on one surface to the ther, and thus restore the equilibrium. Gravitation operating upon the balls as men the pendulum of a clock, evidently co-operates to sustain their vibrations.

87. The length of time that the bells may be kept ringing by the vibration of the alls, when the air is dry, may excite surprise at first; but it should be recollected but the balls can only carry off at each stroke a portion of the electric fluid, which to the whole quantity in the coating, as the superficies of the ball is to the sur-

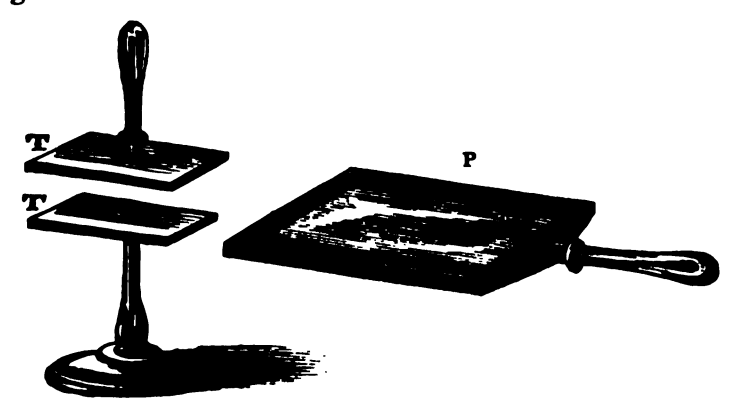
ace of the coating; or less, probably, than as one to ten thousand.

# I the part performed by the Coatings in the case of the Leyden Jar, or Coated Pane.

88. Metallic coatings, employed as in the preceding experiments, are of use in conveying the charge, but do not contain it. The effect of the tin foil is simply to ause the speedy and equal distribution of the electricity wer the surface of the glass, which, being an electric, annot by itself convey the excitement from one part of its surface to another. Hence a pane without coatings can only be partially charged or discharged at one contact.

## Pane with Moveable Coatings.

89. T T represents metallic sheets, which are used as coatings to a mane, also represented in the figure. One of the sheets is insulated by meing supported upon a stand with a glass leg, the other by being held by glass handle.



#### Experimental Illustration.

90. A glass pane, P, held by means of an insulating handle of the same material, is made to touch the knob of an excited conductor by electrical machine on one side; while another metallic kind, passes cating with the other conductor of the machine, is made to touch to me on the other side, in the part opposite to the first mentioned ka. . varying the situation of the knobs, the pane is charged wherever a confaces have been sufficiently in the vicinity of the knobs. If, where the prepared, it be supported by its landle, and one hand of the approximated to one side, while the other hand approximates the cart can only be gradually discharged, as it was charged; the knocks in made to assume, successively, the various positions relatively to the parpreviously occupied by the knobs. But the pane being again charge by the knobs, with the aid of the coatings, T, T, properly applied, the seam are thoroughly and instantaneously discharged, if a conducting comcation be made between the coatings, by means of the hands, arm, at person of the operator, or by any other conductor.

91. The coatings being applied to the pane whilst charging, not an removed, the discharge can only be effected gradually, as the knotes to discharger is made to pass over the charged surfaces, the one on one

the other upon the other side of the pane-

92. Any other conducting substance which will accommodate at the surface of the glass, may be substituted for tin foil. But math a preferable, as they are pre-eminently the best conductors.

Glass Vessel so situated as that Water is made to perform the now of as Metallic Coatings in the Leyden Phial, illustrating the orient Experiment of Cuneus and Mushenbroeck.



quantity of water, of which the exface should not be less than two inches below the brim, and the the the exterior surface should be extend by water to the same height is the inner surface, may, as in the onebrated experiment of Coners and Mushenbrocek, be charged and dicharged by the same means as the pane or plual coated with the folthough less advantageously.

94. Gold, stater, or copper last meta he filings, or mercury, has is substituted for the contings of a Leyden jar. When metallic ates are glied to the surfaces of a peac or jar within the space must's ablotted to the tin foil coatings, the

discontinuity of the conducting surfaces causes the passage of the electrics; from one portion of the surface to another to be indicated by splended corruscations, which will hereafter be fully illustrated.

## Imperimental Demonstration that the Charge of a Leyden Jar does not reside in the Coatings.

25. That the charge does not reside in the coatings, may be proved by assoving them, touching them with the hand while separated from the lase, and afterwards replacing them, and making a communication between them. A discharge will take place in the same way as if they had not been removed.



96. Instead of tin foil coatings, a metallic case is made just large enough to receive a tumbler with ease, and reaching about two-thirds of its height. A hollow cylinder of the same material is made so as to fill the cavity of the tumbler to the same height as the case reaches on the outside, and yet so loose as to be removed without difficulty. The tumbler being charged in the usual way, the cylinder may in the first place be lifted out of the tumbler by means of a glass rod, and the tumbler in the next place grasped at the brim, and lifted out of the case without destroying the charge. This is rendered evident by reinstating the tumbler in its case and the cylinder in the tumbler; and by means of the discharger making a conducting communication between the case and the ball of the wire communicating with the cylin-

der. An electrical spark will then pass with the usual noise. Or, if the circuit be established by touching the knob with one hand, and the case

with the other, a shock will be experienced.

## Of Electrical Batteries.

97. A series of coated jars being placed side by side in a box, and all the inner coatings being made to communicate with each other, and with a ball of metal, by means of metallic rods; and all the outer coatings being made to communicate with each other, and with another metallic ball, by strips of tin foil, the jars thus associated are

called an electrical battery.

98. Charging and discharging an electrical battery, however extensive, is just as simple, and is performed in precisely the same way, as in the case of a single jar. To charge a single jar, or a battery, the different coatings must be made to communicate, severally, with the different conductors of an electrical machine, in operation, either directly, or indirectly through the floor of the apartment or other conducting medium.

99. To effect a discharge, either one or several conductors must be made to form a circuit from one coast to the other, either unbroken, or if interrupted, the user val, or the sum of the intervals, must not exceed a certain distance, called the striking distance, and which vance was the power of the electrical machine.

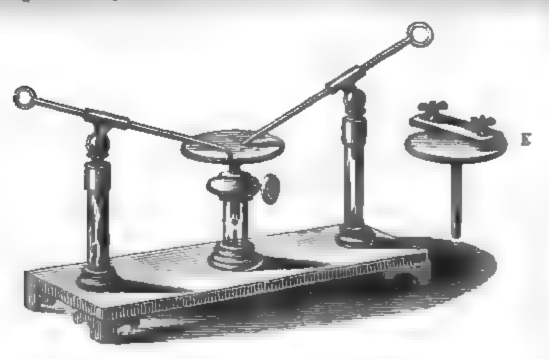
according to the number of jars to be supplied, and a quantity of the electric fluid generated by the mach is But the discharge appears as quick from a great number as from one, notwithstanding the numerous ramificance through which the electricity has to pass.

## Electrical Battery of thirty-two Jars, each thirteen inches in hage, of five inches in diameter.

of the canopy over the hearth of my lecture room, in the vicinity of plange electrical machine. With the positive conductor of this, it is of easily made to communicate by a metallic rod. From the outer couldn't wire is extended to one of the iron columns of my lecture room, along the it afterwards descends to the table, and, when the battery is in use, at sheet of metal on which the battery discharger is placed.

#### Description of Henlay's Universal Discharger.

102. This instrument has been employed to facilitate the exposure of being the discharge from a Leyden jar, or battery.



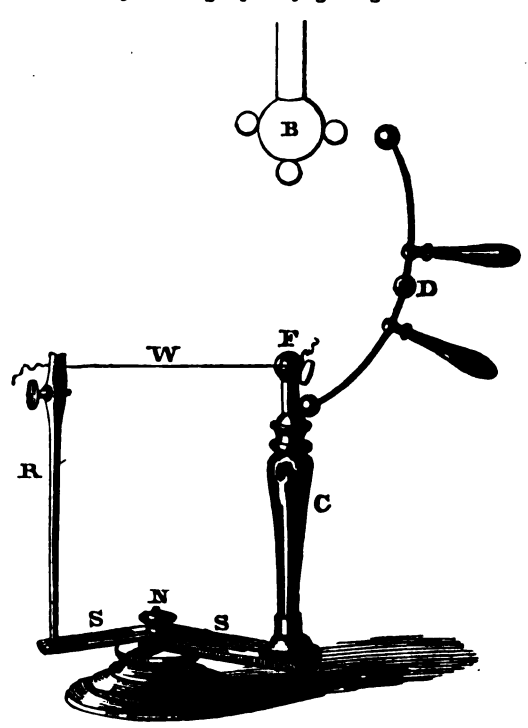
103. It may be understood from inspection. Two rods are supported upon pillars, to which they are secured by universal joints having not only liberty to not on a pivot in almost any direction, but also to be slid through a spring secket are to lengthen or shorten the portion of the rod between the socket and the point of the wires are pointed, but near the points are screws out so as to finish to brass balls which may be screwed on or off.

104. Between the columns is a little stand, which may be altered in height by means of a set screw. Upon this stand, an object to be made the medium of a discharge may be placed, the ends of the wires being in due contact with it. To one of the eyes at the other end of the rods, a wire or chain, communicating with a contact of the battery, may be affixed. By means of the common discharger (c. 76) a communication being then made between the other rod and the other coating of the battery, the circuit is completed, and the whole charge of the battery passes through the body upon the stand.

105. In some cases where pressure is required, the stand represented at E is employed. Thus some gold leaf, being compressed between two very small panes of these, kept together by the screws with which this last mentioned stand is furnished,

be deflagrated, and incorporated with the glass.

#### Battery Discharger for deflagrating Wires.

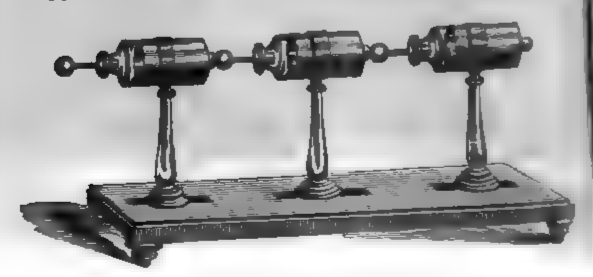


106. This apparatus is employed by me in lieu of Henley's universal discharger above described, being better adapted to my apparatus and mode of operating. Two brase plates are secured to the pedestal by a screw bolt, N, which passes through a hole made in each, near one extremity; the plates are thus allowed a circular motion about the bolt so as to be set in one straight line, or at any angle with each other. On one of the plates near the extremity not secured by the bolt, a brass socket is soldered, into which a glass column is cemented, surmounted by a forceps. At the corresponding end of the other plate there is a brass rod, R, perpendicular to the plate and parallel to the glass column. This rod is also furnished with a forceps. Between this forceps and that at F, supported and insulated by the glass column. C, a wire is stretched, which may be of various lengths, according to the angle which the plates, S, S, make with each other. The pedestal should be metallic, or have a metallic plate at bottom, in communication with the external coating of the battery. This being accomplished, it is only necessary to charge the battery, without subsequently breaking the communication between the inner coatings of the jars and the prime conductor by which the charge is conveyed. In that case, touching the conductor is equivalent to a contact with the inner coatings of the praction as electrical results are concerned. Hence, by causing one of the know of the discharger, D, with glass handles, to be in contact with the insulated foreign f mitthen approximating the other knob to the prime conductor, B, the charge of the tery will pass through the wire, W, as it cannot descend by the glass original reach the operator through the glass handles.

## Of Electrical Excitement by Induction.

phial, or coated pane, already illustrated, where one seface being in contact with an excited conductor, a tendecy is induced in the electricity on the other side of the electric to leave it. Agreeably to the Franklinian them this arises from the self-repellent power which is presumed to exist between the particles of the electrical fluid.

Apparatus for the Illustration of Electrical Induction.

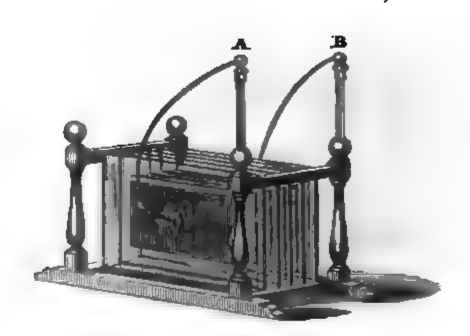


108. If the outer and inner coatings of two or three usual ted jars be made to communicate, and the coatings of each extremity of the series be brought into communication with the conductors of a machine in operation. It usual when one jar is to be charged, it will be found that a charge is received by all; and in discharging them a spark may be perceived to pass between each jar, if a small interval be left. The effect of the discharge is less than that which would be produced by means of one part in this case the surfaces are said to be charged by induction.

109. The number of jars which can be thus affected a greater or less, according to the intensity of the electroity evolved by the machine, and the aggregate thickness of glass interposed.

#### A new Apparatus for the Illustration of Electrical Induction.

110. As the prevailing theories of electricity cannot be understood without
 correct idea of electrical induction, I have constructed, for the purpose of rendering it more intelligible, the apparatus described in the following article. The surfaces oppositely charged, being in the case of panes exactly mike, renders their commutable relation more easy to understand; and the process, as it proceeds in them, having a greater resemblance to that ascribed to voltaic series, may hereafter be more advantageously cited as a smeans of illustration.



about two inches from the edges, are situated in a frame at the distance of about two inches apart. A metallic communication is established between the inner coating of the first pane in the row, and that of the second pane immediately opposite, by means of a spiral spring of wire, which, by its pressure, keeps its place, and produces a close contact between its termination and the tin foil. A similar spring is interposed between each pair of coatings. The external coating of the first, and that of the last pane in the series communicate severally by wires with metallic knobs, A, B, supported upon and of course insulated by glass pillars; the coating of the first pane communicating with the knob at A, that of the last with the knob at B.

112. This apparatus, like a Leyden jar, may be charged in either of three modes. That in which only the positive pole of the machine employed is insulated from the earth; that in which only the corresponding negative

pole is insulated; and that in which both poles are insulated.

113. When the operation is performed with the positive pole insulated, the negative pole communicating with the earth, the surcharge induced in the coated surface of the first pane expels from the inner coated surface of that pane a portion of electricity, which is of course driven through the spiral into the nearest coated surface of the pane next in order. The surcharge induced thus in the nearer surface of the second pane, causes the other surface of this pane to give up electricity to the nearer surface of the third pane: so that by a repetition of the process, every pane will be charged; the last surface or the external surface of the fifth pane, relinquishing a portion of its electricity to the earth.

114. During the access of the charge, the apparatus only receive in a cession of the fluid on the external coated surface of the first pane, and an a portion from that of the last pane. In the other surfaces the quarter not altered; since whatever one loses is gained by another with when a communicates.

of insulation, it will be the converse of that above described. Electrical being abstracted from the external surface of the fifth pane, used being accumulated upon that of the first pane, the internal surface is fifth pane becomes positively excited at the expense of the nearest state the fourth pane, which of course becomes negative on the surface is robbed, and positive on the other side, at the expense of the coated state of the third pane. Thus by a successive inductive influence, transfer from pane to pane, every other surface is negatively charged, causing the which alternate with them to be charged in the opposite way.

poles insulated, the two processes above described co-operate simultaneous since, while electricity is abstracted from the external surface in direct communication with B, on the right, it is accumulated upon the external surface communicating with A, on the left; so that, by the inductive process estimated in the communication with B.

pane becomes charged.

of all the surfaces may be effected by making a conducting communication between the external surfaces of the first and fifth pane by touchast knobs A and B, severally communicating with them, by a discharge. It only difference consequent to the different modes of charging is in the control of the charge relatively to that of the earth, which is conventionally the

fied with neutrality.

116. Each mode of charging is productive of a different state of the pass relatively to the neutral state. Whichever of the knobs A, B, and of course the surfaces of the first and fifth pane communicating with them is to case munication simultaneously with the earth and a conductor of the nather, must have the same state as the earth, and must consequently be much-The other knob, and its connected conted surface, must have the exchange of this conductor. It follows, that in the case in which one knob, A. ibout communicate with the excited positive conductor while the knob. B. com municates with the other conductor and with the earth, the negative exfaces of all the panes, excepting the external surface of the lifth pant, wi be positive relatively to the earth. And, vice versa, that if A commonant with the negative conductor, B, being uninsulated, as above described, id the positive surfaces of the pane, excepting the first, communicating with A. will be negative relatively to the earth. When both conductors of the nechine are insulated, and severally while excited in communication, the or with A, the other with B, the coatings of the central pane will be negral. while all those between that pane and the knob communicating with the positive conductor was be relatively positive, while all on the other sair of a will be relatively negative. To the student, who shall follow me in this etplanation, it will be quite evident that positive and negative electricity and relative states, and that a body, E, may be negative as respects D, yet postive as respects F. When this apparetus is discharged as above described the surcharge in the surface of one of the extreme panes, comprasing the deficiency in that of the other extreme pane; and, at the same time, the equilibrium is restored between all the intermediate surfaces. As repects these there has been no loss or gain of the electric fluid. The actual changes have only amounted to a transfer and retransfer from one set of the surfaces to another.

## Exposition of the difference between Induction and Conduction.

119. There is a great analogy between the process by which an excess electricity in one body is conveyed to another through a conductor, and hat in which the air condensed in one vessel is conveyed to another through tube, in order to attain an equilibrium. But when the process by which L body, surcharged with electricity, electrifies another body, inductively, is compared with any which can exist between a cavity surcharged with air. and another in its vicinity, no analogy can be discovered. However air may be condensed in one receiver, it has no influence upon the quantity which can be retained by a neighbouring receiver with which no channel of communication exists. That removal of all obstruction to the transfer I the redundant fluid, which is the effect of the conducting process, would fatal to the process of induction. This requires that no means of conreyance shall exist between the electrified body and that subjected to its inluctive influence. Moreover, the excitement produced by conduction, is the same as that of the exciting body, while that which results from induction s the opposite of the kind by which it is induced. Nevertheless, a body sharged by conduction may produce a charge by induction; and a charge y induction, may be discharged by conduction.

## Of Dynamic Electrical Induction.

120. Besides that kind of statical electrical induction, which I have been endeavouring to explain, there is another kind which may be called dynamic electrical induction. For our knowledge of this species of electrical reaction, we are indebted to Faraday. It is in fact to this habitude of electricity, that the Faradian currents are attributable, respecting which I have treated briefly in my Treatise on Electro-magnetism, page 60. The experiments of Henry, mentioned page 57, of same treatise, afford an illustration of the analogy between the dynamic inductive influence of electricity proper, and that excited by galvano-chemical reaction.

#### MEANS OF DETECTING OR MEASURING ELECTRICITY.

121. It has been seen that the property which light bodies have of separating from, or approaching to, each other when electrified, has been of use in showing the nature and extent of electrical excitement.

122. A ball of pith, supported by a radius, suspended from a pivot, so as to be capable of describing an arc of ninety degrees, over a corresponding curved scale, constitutes Henley's quadrant electrometer, employed in the experimental illustrations. (e. 130.)

123. Bennet's electrometer has been described, in which metallic leaves are suspended within a glass cylinder, to a metallic cap; slips of tin foil

being pasted on the glass, opposite and parallel to the gold leaves.

124. This last mentioned instrument is sometimes more properly called an electroscope; as it is better calculated to discover electricity than to measure it.

125. The efficacy of the gold leaf electroscope is much increased by the addition of two metallic disks, one soldered to the cap, the other attached

to the foot by a hinge; so as that it may be placed parallel, and a the first mentioned disk as it can be without touching. In the capacity for electricity of the disk attached to the cap is found a creased by induction; so that it will receive a surcharge. When the are separated, the excess of electricity received while they were mother, is indicated by the divergence of the leaves.

126. The instrument thus constituted, is called the condensing

meter, of which an engraving and description is annexed.

#### Description of the Condensing Electrometer.

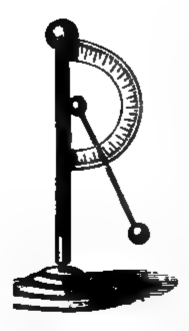


127. The condensing dis ter, of which the annexed in representation, differs from t nary instrument, in being fa with two metallic disks, out ed to the canopy, C, the of held by a wire. The wind nates in a hinge at A, by disk which it supports mey b to approach, or retire from th disk. The metallic hinge ( nicates, by a strip of tin in other strips of the same which are pasted on the gl already described in the case gold leaf electrometer. (c. 51

128. In order to put this instrument into operation, the disks a quite parallel, and as near each other as possible without contact on touching the cap with an electrified mass, containing a charge tricity otherwise too low to affect the leaves, and afterwards remove disk D to the distance of two or three inches, the leaves diverge. A gence of the leaves of the condensing electrometer may be prosupporting a zinc disk of about six inches in diameter, in the h otherwise, so as to have a communication, directly or indirectly, w pedestal of the electrometer, and placing on it, from ten to twenty a disk of copper of the same size, held by a glass handle, and at a moval bringing the copper disk in contact with the cap of the che ter. By these means a charge is imparted to the cap, which, wh outer disk is removed, is evinced by the divergency of the leaves our knowledge of the last mentioned method of producing electric the contact of heterogeneous metals, we are indebted, I believe, to the brated Volta.

129. I have constructed an electroscope, with a single leaf, to wibrass ball may be approximated by a micrometer screw. This is sensitive than any electrometer which I have seen on the usual When furnished with a cap of zinc, if a plate of copper be placed cap, and then lifted, the leaf will strike the ball. This instrument act as an electroscope and as an electrometer, as it detects and measure minutest degree of excitement.

#### Description of Henley's Quadrant Electrometer.



130. Henley's electrometer consists of a little wooden column, supporting a semicircle of ivory, or of wood covered with white paper, graduated near the periphery into 180 degrees. At the centre of the semicircle there is a pin, from which a moveable radius terminated by a pith ball is suspended. This radius is sufficiently long to allow the ball to reach to the base of the column, against which, when left to itself, it But when the ball and column are electrified, the ball moves off from the column, together with the radius to which it is affixed. But the radius being secured to the pivot at its upper end, the ball must describe a greater or less portion of a circle, which is at the same time indicated and measured by the graduation.

ription of the Single Leaf Electrometer, by which the Electricity, excited by the touch of heterogeneous Metals, is rendered obvious after a single contact.

II. A single gold leaf is suspended from a disk of zinc six inches in diameter, sh constitutes the cap of the instrument. Opposite to this single leaf a ball is sorted, which may be made to approach the leaf, or recede from it, by means of a screw. Of the same size as the disk which forms

the cap there is a copper disk with a glass handle,



accompanying the instrument."

132. The electricity produced by the contact of copper and zinc, is rendered sensible in the following manner. Place the disk of copper on the disk of zinc, which forms the cap of the electrometer. Take the micrometer screw in one hand, touch the copper disk with the other, and then lift this disk from the zine. Usually, as soon as the separation is effected, the gold leaf will strike the ball, if the one be not more than the twentieth of an inch apart from the other. Ten contacts of the same disks of copper and zinc will be found necessary to produce a sensible divergency in the leaves of the condensing electrometer. That the phenomenon arises from the dissimilarity of the metals, is easily shown by repeating the experiment with a zinc disk, in hen of a disk of copper The separation of the homogeneous disks will not be found to produce any contact between the leaf and ball. I believe this to be the only

le in which the electrical excitement, produced by the contact of heterogeneous

tals, can be made evident without the aid of a condenser.

33. It is probable that the sensibility of this instrument is dependent on that perty of electricity which causes any surcharge of it, which may be created in a decting surface, to seek an exit at the most projecting termination, or point, nected with the surface. This disposition is no doubt rendered greater by the ximity of the ball, which increases the capacity of the gold leaf to receive the charge, in the same manner as the uninsulated disk of a condenser, (c. 27,) inmees the electrical capacity of the insulated disk in its neighbourhood.

34 It must not be expected that the phenomenon above described can be pro-sed in weather unfavourable to electricity. Under favourable circumstances, 1

For the experiment with this electrometer, a metallic handle would answer. being of glass, enabled me to compare the indication, thus obtained by my insment, with that obtained by a condenser.

have produced it by means of a smaller electrometer, of which the data to me two and a half inches in diameter

135. The construction, as respects the leaf and ball, regulated by the more crew, remaining the same, the cap of a condensing electrometer and its thin my be substituted for the zine disk.

#### Description of Coulomb's Electrometer.

186. The electrometer of Coulomb is suitable rather for the incurred than for the illustration of electrical phenomena. Yet, as it may become to convey an idea of the principle of this instrument, I shall quote -: > Treatise on Heat and Electricity of the distinguished Dr. Thomson .... gow, a description, accompanied by an engraving of the clockment ! question in the most simple form. Alluding to the gold leaf circumst (e. 54,) or that in which straws are used in place of gold leave. Thomson observes:-

137. " In these and many other common electrometers which I think it make to describe, the instrument cannot be considered as a true measure of the seed of electricity; because as the two straws or the two slips of gold lear separate ther and further from each other, it is evident that gravitation will us not more powerfully to bring them back again to their naturally vertical position. the repulsive force of the straws, or leaf, is not proportional to the distance to well they separate from each other. These instruments cannot of course be expense. to measure the energy of electricity.

138 " But the electrometer of Coulomb is free from the company of the coulomb is free from the coulomb is f

It is represented to the margin. It consists of a glass vesses in a lid also of glass, in the centre of which a must hole is Through this hole passes an untwisted raw silk thread har solling, and fixed at the top to a micrometer, by means of waste may be turned round any number of degrees at pleasure 1. silk thread is attached a very fine gum lac thread, H. harate each extremity a small knob. This lac needle, with a tem weighs only one fourth of a grain. A small hole is called a small hole is called a side of the vessel, at A, through which passes a fine way an excited below the placed in contact with the knob at A, the knob at the other tremity will acquire the same electricity as the excited body. This electrons are communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the communicate to the knob of the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk thread with the lac needle suspended by the milk

previously almost in contact, and the two knobs wil, repel each other. The more ble knob attached by the silk thread will separate from the other, and the question of electricity will be proportional to the distance to which it is driven off

139 "Coulomb's electrical balance is an instrument intended to means in

quantity of electricity in bodies, and indispensable in accurate experiments.

140 It should be understood that in this instrument, the knob of the expense. needle may be made to resist sufficiently its removal from that supported by wire, by twisting the silken fibre. Coulomb contrived a more perfect and come cated electrometer, upon the same principle as the one which I have described " furnished with graduated circles for measuring toe distance between the balls, the extent of the torsion given to the suspending filament t

141. By means of this apparatus, Coulciob contributed an observation provide made by the Earl of Stanbepe, that the density of electricity in the electrica in sphere, surrounding an excited body, during he inversely as the square of the tance from the charged body. Concombiniferrid, from the law thus assumed to state and from ingenious and accurate experiments tending to corroborate his interest that the electricity accumulated about a conducting body is entirely superious, none of it existing in the interior of the body. He also, as Dr. Thomson concerns. " proved by very simple but convincing experiments, that electricity depostes and upon bodies according to their surfaces, that it has no more attraction for one bely than for another also, that if two bodies, having the same surface be placed # contact, whatever their nature may be, any electrical surcharge in either will #

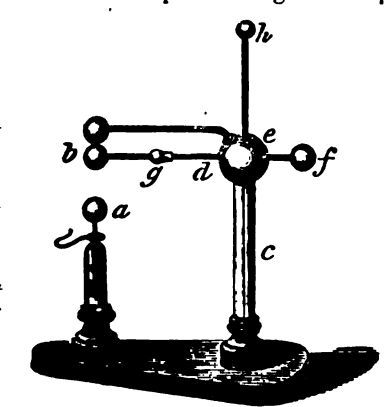
<sup>\*</sup> I think I have seen an effect from a disk only an inch in diameter or from t zine disk having a handle with a copper socket

t The description of this last mentioned instrument, with engravings, occupthree pages in the Treatise of Dr. Thomson, alluded to above, which is more specified than I deem it expedient to devote to the same purpose.

**livided** equally between them." Allowance is to be made for the obstruction arisme from the diversity of conducting power, which, however, only delays the equal**zation**, but does not prevent it from taking place.

### Of Cuthbertson's Balance Electrometer.

142. The following figure affords a faithful idea of this instrument. which I shall proceed to give an explanation.



143. The brass ball, a, surmounts a rod of the same metal, upon the lower end of which a cork is secured. This cork may be slid upwards or downwards, within the glass tube supporting it, like the piston of a syringe. By these means the ball, while insulated by the glass, is liable to be so varied in its height, as to be supported at any eligible distance from the ball, b, beneath which it is situated. This last mentioned ball, b, and the rest of the apparatus with which it is associated, is supported and insulated by means of the glass leg, c. The ball, b, is attached to the end of the rod,

d, which is balanced upon a fulcrum within the hollow globe, e, in the same way as the beam of a steelyard. The ball, f, is made just heavy enough to counterbalance the ball, b, together with the sliding weight, g, when this

weight is at the greatest possible distance from the fulcrum.

144. The globe, c, supports two rods, one proceeding from it vertically and surmounted by the ball, h, the other proceeding from it horizonally, so as to be in contact with the ball, b, as represented.

145. When, previously to the charging of a Leyden jar, the ball, a, is made to communicate with one coating, the outer one, for instance, while the rest of the balls and the lever communicate with the other coating, the balls, a and b, are made to attract each other with a force proportionable to the intensity of the charge and their proximity. Hence, by approximating a to b, or by sliding the weight, g, further from the fulcrum, the resistance of the counterweight, f, may be rendered inadequate to counteract the electrical attraction produced by a very low charge. Whenever this takes place, the ball, b, will descend until the interval between it and a, is so small, that the electric fluid can leap through it so as to discharge the jar or battery. On the other hand, by increasing the distance between a and b, or lessening that between the fulcrum and g, a proportionably higher charge will be requisite to overcome the resistance of the loaded ball, f, so as to allow b to descend sufficiently to enable the discharge to ensue.

146. It follows from the construction of this electrometer, as explained, that we can cause successive discharges of exactly the same strength, by keeping the distance between the balls a and b, and that between the weight g, and the fulcrum unchanged. It must likewise be evident, that the intensity of the charges, and consequently of the discharges and shocks given, may be regulated by altering the distance between the balls, or that be-

tween the sliding weight g, and its fulcrum within the globe, e.

147. There is a most interesting analogy between the descent of the ball b, and that of the cloud of an incipient tornado. If some pith balls be supported in a shallow cup, upon the ball a, they will be seen to ascend towards the ball, b, when descending in obedience to electrical attraction, m in case of meteors of the kind abovementioned, water and other must bodies may be seen to rise up toward the descending cloud.

#### EFFECTS OF ELECTRICITY.

148. The separation or approximation of electrified dies, the extrication of light and heat, and the shock in to the animal frame, having been all, more or less, support of discussion, or adduced as the means of experimental illustration, it may now be proper to display a greater riety of electrical phenomena, and such as being a complicated require undivided attention on the part those who would comprehend them.

Of Electrical Attraction.—Of Electrical Light.—Of E trical Ignition.—Of the Electrical Shock.—Of Mitigat the Effects of Electricity.

## Of Electrical Attraction.

149. Under this head are placed both the separated and approximation of light bodies, when electrified; in the former, though commonly ascribed to repulsion really, as I conceive, the effect of attraction.

Revolution of a Sun, Planet and Satellite.

Fig. 1.

150. A hollow brass globs, 1, is rendered much heaver on side by running into it a quar of molten lead, sufficient to occ about one-third of the cavity. these means, when supported o pivot, it preserves a proper post although on the other side not nished with lead, it is made to port an arm and two balls. larger, representing a planet, other smaller, representing its s lite. These are carried upon different ends of a wire pass through their axis, and balas upon the point of the arm, so t the balls may counterpoise e other. From the larger arm, from the smaller ball, points P. project, as represented in Fig. 2.

B P A

Fig. 2,

51. When the sliding rod of the conductor, R, is brought sufficiently to the central globe of this apparatus, the machine being in operation, ensues a complicated revolution. The planet and its satellite, attached e arm B, are balanced upon a pivot, formed of the recurved and pointed ination of the larger arm, A; meanwhile the pivot revolving as the S, turns upon its axis, carries the planet and satellite in their orbit, e they revolve about the pivot as their common centre of gravity.

i2. According to the Franklinian theory, the above described rotatory one result from the attraction of the surrounding medium, operating essively at every part of the orbit, to separate the electrified air from

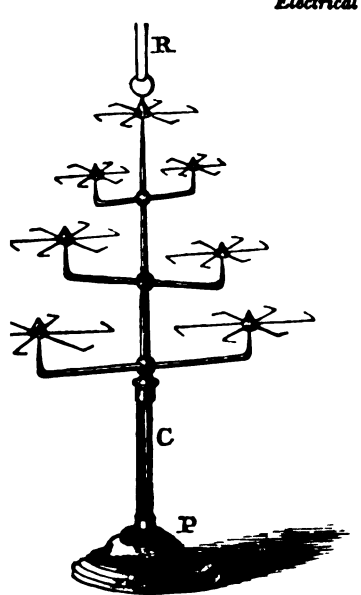
netallic points by which it is electrified.

i3. It must be admitted that as these rotatory motions may be produced, her the excitement be negative or positive, the rationale is more diffiof conception upon the theory of one fluid than upon that of two. It seessary, agreeably to the Franklinian doctrine, to ascribe all cases in h electrified masses separate from each other, to attraction between and the matter of the adjoining medium. This process becomes exely difficult to follow in the "mind's eye," when the masses thus sepal are undergoing a rapid change of position.

i4. The explanation, supposing two fluids to exist, is simply that bodies arly excited repel each other; and consequently that repulsion arises sen the points and the air which they electrify, whether the excite-

be vitreous or resinous.

#### Electrical Tree.



155. I believe that an apparatus resembling that of which this figure is an engraving, is named in Pixii's catalogue, "Arbre Electrique," a name which I have used for want of a better.

156. It consists of several sets of branches formed of wire. Each set is associated by a common hollow brass cone, into the apex of which a recurved wire, forming a principal branch of the electrical tree, is so introduced as to form a support and a pivot, upon which the cone and its branches may rotate. Each rotatory branch is recurved, and terminates in a point; the points in each set projecting in the same direction, so as to co-operate in producing a circular motion.

157. The branches are put into operation by communicating with the machine, as usual, by the rod R. The excitement thus received cannot pass off by the trunk C, which is of glass, cemented into an upright brass rod above, and the pedestal P, below. It can hardly be necessary to add, that the ra-

e of the rapid rotation of each set of the branches is analogous to that of the ding experiment. In consequence of their being similarly surcharged with icity, or similarly deficient, the adjoining neutral medium attracts the air and anches apart with energy, and thus causes them to recede from each other as as they come into proximity.

. To render the Franklinian rationale more intelligible, as applied to this examt, and that of the minature sun, earth, and moon, it may be observed, that

the excitement of the machine being communicated by the cliding red it, to central ball, and of course to all of the metallic wires there with associated the fused through the points to the air in their vicinity. Consequently the points (fig. 2.) and the air electrified by them, being similarly succlarged or much ficient, must be attracted by the adjoining neutral medicine, and not attracted other, they are made to separate rapidly, or to move in opposite direction. I process being resterated with inconceivable speed, the revolutions present a proportional velocity.

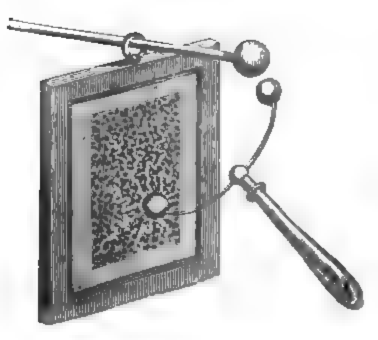
Electrical Hail.



159. This experiment, while Pixu's Catalogue described as a Electrique, 's electrical had an another illustration of the possible which may be produced a light dies by electrical attractive tallie rod supports one bill on the bell glass, another without m to be in contact with the ball another rod, R, proceeding famile conductor of the large electrical chine in operation. The large being by these mosne intrody trified, attracts some of the public which he upon the metallica which the bell is situated and in should communicate with the tons of the martune Assemble pith balls come in contact was excited agreeably to the ground they recede from each other, and attracted by the oppositely cannot dish. Reaching the dash they di the same electrical state as at f and of course are liable to be attened again Meanwhile other hill undergoing the same routice proing that contrariety of merian which characterizes the full of the

#### OF ELECTRICAL LIGHT.

Experimental Mustration of the efflux and afflux of the Electric Fluduring the charging and discharging of coated surfaces, as render evident by means of discontinuous coatings of Metallic Filings.



discharging of a conpane has already be illustrated and explane. The process is, however rendered much more a teresting, when, into of a continuous contaof foil, a covering of matallic filings is applied, as to leave a multitude a minute intervals between the particles of the meait is only necessary thave the discontinuity of raring on one of the surfaces; the other may be coated with tin foil as sai.

161. The tin foil coating should, by means of a strip of the same submee, communicate with the ring attached to the wooden frame in which pane is secured. Let the pane, thus prepared, be suspended by the g, as represented in the engraving, from a rod affixed to the conductor a powerful electrical machine in operation. As represented in the figure, a discharger be so held, in contact with the surface coated with filings, to carry off electricity from it, allowing the other surface of the pane to proportionably surcharged. In the next place, let the discharger be so anted, as to complete the circuit between the surfaces, allowing the sur-



 $\mathbf{R}$ 

charge in the one to rush to the other. By these means the efflux and afflux of the electric matter will be indicated by coruscations of electric light with an indescribable splendour, which will appear so long as the surface coated with tin foil remains in communication with a machine sufficiently active, and the situations of the discharger are alternated as above described.

162. Analogous results may be obtained by means of a jar represented by the preceding figure, which has a hook wherewith to suspend it; or still more advantageously by means of a large carboy, silvered with an appropriate amalgam within, and on the outside furnished with a discontinu-

ous coating of filings.
163. If the discharger employed in this experiment have a glass handle, either the metallic socket, S, into which the handle is cemented, must be touched by one of the fingers, or a wire must be at-tached to it, making a communication with the negative conductor, directly or indirectly. Otherwise, the external costing being insulated, electricity could not escape from it, and of course the inner surface could not be charged as already demonstrated.

164. The adjoining cut represents a glass carboy of about five gallons in capacity, coated internally by means of the amalgam usually employed for the purpose, externally by brase filings, as in the cases of the pane and phial above described. It is situated under the projecting ball of the prime conductor, so that the knob at the top of the rod, pro-ceeding through a cork from the internal conting, may be in contact with the slid-ing rod, R, of that conductor. The wire, W, is supported on a pedestal in contact

with the external coating of the carboy, and in communication with the conductor of the machine. Hence, when by the operation of the electronic the internal surface of the glass is becoming charged, the exape of the from the external surface is indicated by cormications of light, and counds, as in the preceding experiments, and at the instant when the quiros sufficient intensity to jump through the interval between the hind to the ball, and that supported by the wire, at K, the deficit created ternal surface being restored at once, the cormications and spatiting are a splendid.

Long Zigzag or Erratic Spark, contrasted with the Sha Straight Spark.



represent the different forms and of the electric spark, which also tween a large and a small ball, ingly as they are made negative tive. The long and zagzag, a spark A, takes place between a mattached to the positive pole, and one associated with the negation one associated with the negation of course only be obtauted and can of course only be obtauted succession.

166. In no respect do the plat of mechanical electricity appear ( voorable to the Franklinian that more inexplicable according to the of two fluids, than in the diversit electrical spark in passing between and a large metallic ball, according manner in which the balls are 🖦 with the positive or negative pole machine. When the small ball as to the positive pole, the spark 1 comparatively narrow, and of a shape, such as lightning is oficaassume; but when the situation bals is reversed, the spark is strag thick, not one-third as long, and t of a zigzag shape can be observed

167. According to the Franklinian theory, when any body is highly charged with electricity than the adjoining bodies, the excess fluid is attracted by them, while it is inadequately repelled by the a quantity of the electric fluid, with which they are imbued. It follow when a small globe is made positive in the neighbourhood of a large the excess of electric matter in the former is attracted by all the neglective metal in the latter. When the small globe is made negative metal of which it consists attracts all the electric matter in the large Hence there is this difference in the two cases; the small globe being tive, a comparatively small moveable mass of electric matter is attracted at the electric matter is attracted at the electric matter is attracted.

e, a large immoveable mass of metal: the small globe being made negative, a large moveable mass of electric matter is attracted by a small impoveable mass of metal. The charge being in both cases the effect of the mass machine, the attractive power must be as great in one case as in the bear. The forces by which the masses are actuated being therefore equal, is quite reasonable that the greatest projectile power should be attained, bear the small mass is moveable. In that case, it will require less air to removed in order to effect a passage.

168. There is an analogy between the difference which I suppose to tist in the case under consideration, and that which may be observed between the penetrating power of a rod which is blunt, and one which is

pinted.

169. It remains to be shown why a large mass of electric matter will discharged in a spark when there is sufficient proximity; although that extric matter be situated in the large globe, and attracted by the other, and attracted by the other, as above stated, it would not pass without

mt proximity.

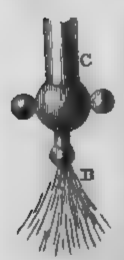
170. It must be evident that attraction increases as the distance between bodies which exercise it lessens. Of course the attraction of the small lobe must always act more powerfully on those portions of the electric mid, which occupy the nearest parts of the positively excited globe. But is difference of distance, and consequent diversity of attraction, increases a the globes are approximated. Thus that portion of the electric fluid hich sustains this pre-eminent attraction, will be accumulated into a smoid; the acuteness of which, and attraction causing the acuteness, in-reasing with the proximity, there will at last be sufficient projectile and smetrative power to break through the air, and thus open a passage for whole of the quantity attracted by the small negatively excited globe.

171. When, by the process last described, the fluid is made to leap trough a comparatively small interval, by the concentrated attraction excised by a small negative ball upon the expanded surface of electric material diffused through a large globe; the air does not become sufficiently contended to resist it before it reaches its destination, and, of course, it cannot become the erratic form which would arise from repeated changes in its

purse, as in the instance of the long spark.

172. Faraday conceives that his late researches warrant the inference set the diversity between the positive and the negative spark, is affected the nature of the gas in which the experiment is made; and he infers that the difference is due to the influence of the gaseous substance interposed. I does not appear to me that he has sufficiently considered the fact, that he differences between the sparks in the experiments above represented and lescribed, (e. 165, &c.) is the effect simply of the relative position and dimensions of the balls between which they pass. That it is not dependent on he nature of the electricity must be evident from the circumstance, that the long and the short spark were produced by the positive excitement in the experiments adduced by me. (165, &c.)





178. When the machine is in active operation, to prime conductor insulated, the electricity will be a from a small knob attached to it, so as by the material light to exhibit the form of a luminous brud, presented at B, in the adjoining figure. For the parties of this phenomenon, it is necessary that the fluid shall be condensed into a small prominent to as, agreeably to the preceding explanation, to have penetrating power. This it cannot possess, when, to same intensity in the generating power, a large positively electrified. In that case, the electric operation and to broad to produce a passage that surrounding non-conducting air. A small ball, say

electrified, can only be productive of a diffuse attraction for the electricity in the atmospheric medium around it; so that it has less about it any penetrating power, than when acting upon the electricity in paratively large globular conductor, as in the preceding illustration, when the knob is on the negative pole, it may be productive of a lappearance in its immediate vicinity, where the electric matter, entering the adjoining space, becomes sufficiently intense to be producible; but it does not produce the striking appearance of the luminous

174. As, agreeably to Du Fay's theory, the knob, whether varates resinously electrified, is surcharged with an electric fluid, the proposer ought to be as great in the one case as in the other; and the apark and the brush should be producible in either case.

175. It has been mentioned that Faraday has of late endeavoured to that the difference between the forms of the positive and negative a greater or less, according to the gas through which they may be made place. An analogous change is also alleged by him to be produced at tive and negative brushes, which are adeged to vary in their colour, it tive dimensions, with the nature of the gas within which they may be to

## On some Inferences from the Phenomena of the Electric Spark, co in a recent Work on Heat and Electricity.

176. In his valuable work on heat and electricity. Dr. Thomse that if a long spark be taken between two knobs, as when sever tached to the positive and negative conductors of the electrical methe portion of the spark near the positive knob exhibits all the che of positive electricity, while the remaining portion, proceeding frother knob, displays all the characters of negative electricity. All the learned and ingenious author does not state what differences the between the different portions of the spark, and wherefore, if any, excan, without a petitio principil, assume that they are such as to just conclusion, he proceeds to allege that there can be no doubt that spark consists of two electricities; which, issuing severally from a spective knobs, terminate their career by uniting at the non-lumine tion of the spark, which is at a distance from the negative knob of one-third of the interval. Upon these grounds be infers that the petericity occupies two-thirds of the length of the spark, the negative third.

177. I presume, agreeably to the theory which supposes the exist two fluids, that, when the equilibrium between oppositely excited sur

bestored by a discharge, whether in the form of a spark or otherwise, there must be two jets or currents passing each other; the one conveying as such of the resinous as the other does of the vitreous electricity. ourse no part of a spark can be more negative than it is positive, nor more contive than it is negative. Upon this ground, a suggestion of the same author that the diminution of light near the middle of the spark results from be combination of the different fluids at this point, appears to me objectionble; since there is as little ground for supposing the union of the fluids to ake place there as elsewhere. But admitting that the union does take lace as supposed, is this a reason for the observed diminution of light? If, when isolated, either fluid is capable of emitting a brilliant light, should **sot their** co-operation increase the effect?

178. In deflagrating, by voltaic electricity, a wire of uniform thickness, equally refrigerated, the most intense evolution of heat and light is always

midway.

179. In truth, the theory which the learned author sanctions, requires wo postulates so irreconcilable, that unless one be kept out of view, the ther cannot be sustained. It requires that the fluids should exercise an ntense reciprocal attraction adequate to produce chemical affinity, and of course, enter into combination when they meet, and yet rush by each other with inconceivable velocity, not only through the air, but also through the restricted channel afforded by a small wire. If the fluids combine at a point intervening between the surfaces from which they proceed, what besomes of the compound which they form? Is it credible that such a comsound would afford no indication of its existence? But, again, how are wo surfaces, the one previously deprived of a large portion of the negative electricity naturally due to it, the other made as deficient of the positive huid, to regain their natural state? By a combination midway, the resinous und vitreous surcharges might be disposed of, but whence could the vitreous and resinous deficiencies be supplied?

180. Dr. Thomson, in common with the great majority of modern chenists, ascribes chemical affinity to the attraction between the two electriciies combined with ponderable particles. As the combinations between such particles take place only in definite proportions, would it not be consistent that the fluids which give rise to them should combine agreeably to those aws? But if the electrical compound, formed of the vitreous and resinous sloctricities, be decomposable by induction, as the theory in question requires, its constituents must be capable of uniting in every proportion.

181. Agreeably to the late investigations of the celebrated Faraday, equal quantities of the electric fluid are evolved, by analogous chemical changes, from equivalent weights of different ponderable bodies. It may, therefore, be inferred, that in entering into combination, the electric fluid is obedient to

those laws of definite proportion which regulate other substances.

#### Illuminated Globe, Pane, and Tube.

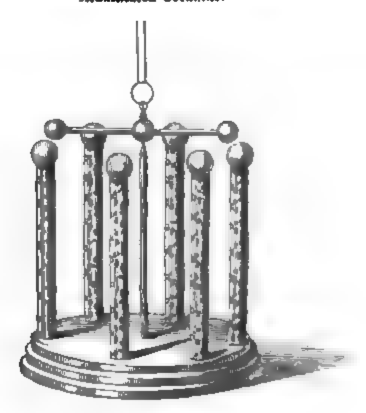
182. Little disks of tin foil are so pasted in succession on a glass globe, pane, or tube, as to leave only a minute interval between them. In consequence of this arrangement, when situated in the circuit between the poles of an electrical machine, the electric fluid, availing itself of the conducting power of the metal, and leaping over each interval, produces as many sparks as there are intervals. At the same time, if these intervals fall within the lines of any drawing, the image representing the drawing will appear at each flash; and when, as in using a large machine, the flashes are almost incessant, the effect upon the eye is nearly as permanent as if the illuminated spaces were inherently luminous.



183. In the case of the globe and tube, represented by figures 1 and 2,1 are arranged in spirals, winding about the globe or tube from one apex or extern. However intricate the route which ingenuity may, by means of the foil, prescribe, it will be pursued by the fluid with fidelity, when the air and dry. Yet the sum of the intervals must not exceed the whole striking 4 the machine, or in other words, the greatest length of its spark.

184. This illustration may be varied by means of apparatus, of which and descriptions will be found in the two following articles.

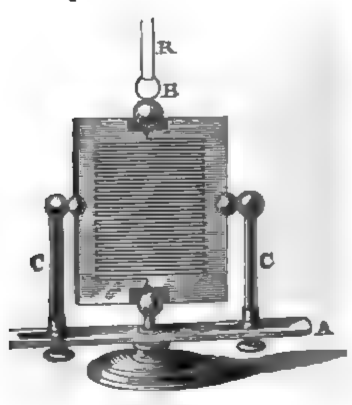
#### Illuminated Columns.



35. A swivel of wire, terminating in knobs, is so balanced upon a pivot, that, in charged, it revolves from the brass knob of the column to which it is nearest, but of the next, and thus successively conveys to each the sparks imparted to it the machine.

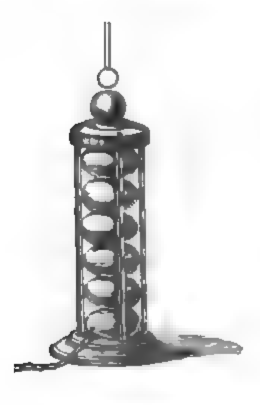
#### Carreux Etincolantes .- Sparkling Panes.

96. This appellation is given in Pixii's Catalogue to the apparatus described in following article.



187. This figure represents a pane, in which the discontinuities in the tin foil attached, as explained, (s. 182,) being made to occur along the lines of a drawing, an image is seen as often as the electric spark passes. The frame, A, having two glass columns surmounted by metallic clasps for securing the pane, is contrived to ac commodate a succession of pance, all of one size exactly, on which different images are delineated, and which may be shown in succession. In operating, the upper ball, B, is situated so as to be in contact with the sliding rod, R, of the prime conductor. At this point, the fluid enters, and coursing horizontally, first to-wards one side, and then towards the other, it finally escapes at the other ball into

foot of the instrument, which should communicate with the cushions of the mane, directly or indirectly.



#### Illuminated Eggs.

188. Between three glass columns a pile of eggs is supported in mutual contact. They are also in contact at top and at bottom with spiral wires; in the one case communicating with a chain, in the other, with a brass knob which surmounts the cap in which the columns terminate, and by which they are secured. The eggs thus situated may be taken into the circuit between the conductors of the machine, or the coatings of a battery, and thus be illuminated by the passage of the electric fluid. The experiment is seen to the best advantage when a powerful machine is employed, as the effect is in that case durable; scarcely any intermission taking place in the flashes. The contents of the eggs appear to be exposed to the eye, as if divested of their shelly covering.

Electric Light in Vacuo.—Aurora Borealis.—Shoating More.



189. The appearances of the electric fluid, when in the act of passing an insulated conductor intensely excited; or during the efflux, or all charging and discharging of coated surfaces, has been sufficiently directed ato.) When an electrical discharge takes place within a cavity from any has been exhausted by an air-pump, the form assumed in much more different to the country of the c the brilliancy of the light is proportionably diminished. Moreover, through which the discharge can take place, is prodigiously greater. It is of an apparatus which the adjoining figure represents, that I am seem exhibit the passage of electricity through an exhausted receiver. After sufficiently removed by the air-pump, it is only requisite to make a comme between the excited prime conductor of the machine (c. 21) and the repasses through the cap, as represented in the figure, in order to have the m tiful coruscations, resembling in miniature those of the aurora boreals confident that this meteor is the effect of electrical discharges, as that arises from that cause. I conceive that such discharges take place not only of the terrestrial poles, rendered electric by frost, into the rare medium er only a few miles above the earth's surface, (98,\*) but also from one par I will venture even upon the bold suggestion, the rare medium to another discharges may take place not only from one planet to another, but even sun, or solar system, to another Since it has been shown by Wheatstone, velocity with which electricity can be transmitted through a perfect connot less than that with which light is projected from the sun, wherefore she unreasonable to conjecture that the former, in order to preserve its equ should pass through distances commensurate with those through which the known to radiate

190 I think it highly probable that the shooting stars of November 13 were caused by the afflux of an immense quantity of electricity to our plus some other planet, or from some remote part of space

191. It may be said that the passage of light is now generally ascribe undulations of an intervening ether. I would in the first place observe remains to be shown that a discharge of electricity is not due to progressive tion of an intervening imponderable fluid ! Besides, if the passage of b

In order that numbers referring to the new edition of the Compendium distinguished from those referring to this treatise, the latter will be precede letter a.

<sup>†</sup> See my Treatise on Electro-magnetism, page 79.

articles, from the most remote of the visible stars, was not deemed an objective Newtonian theory of light, wherefore should there be any difficulty in g a transfer of electric matter to be productive of an aurora or shooting star? I appears to me that the ordinary luminous appearances which we call fall, may be the effect of electrical discharges upon our planet, which become a they enter the terrestrial atmosphere.

## Of Electrical Ignition. Ignition of Cotton by the Electric Spark.



193. Let one of the knobs of a discharger be surrounded by a tuft of carded cotton, containing as much finely powdered rosin as it will retain; and while the other knob touches the outer coating of a Leyden jar properly charged, let the cotton be approximated to the knob proceeding from the inner coating of the jar. A spark will pass, and the cotton will be ignited.

#### Inflammation of Ether by the Electric Spark.

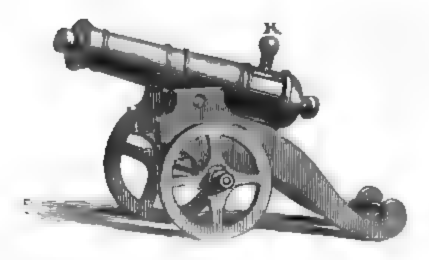
194. This object is conveniently effected by means of a wine glass with a perforated stem, through which a wire passes from a disk of metal cemented to the foot. The wire rises about two-thirds of the height of the glass, terminating in a knob. Up to the middle of the knob, the cavity of the wine glass may be filled with plaster of Paris, or other cement, so as to leave only

a portion of the knob bare. The other should form a thin stratum not entirely covering the apex of the knob.

195. In consequence of its ready vaporization and admixture with air in open vessels, ether is always surrounded by an explosive atmosphere which is readily inflamed by the passage through it of an electric spark, either from the conductor of a machine, or from a Leyden jar. The result is rendered more interesting, when accomplished by a discharge from surfaces electrified by means of water. (s. 93.)

#### Ignition of Hydrogen with Oxygen.





By means of a wire, insulated by being enclosed in a glass tube comented acket, which screws into the touch-hole of the little cannon above represented,

a spark from an excited conductor may be made to cross a small interval when the bore of the piece. Hence, if an explosive gaseous muxture of hydrogen was gen, or atmospheric air be made to occupy the cavity, and be incurred by any mixture will explode on presenting the knob, K, of the insulated was a known conductor of the machine when in operation. It may be fired by the fact of nose of a person supported upon an insulated chair or stool, and community a the same time, with the prime conductor sufficiently excited.

## Of the Electrical Shock.

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197. This effect of the electrical fluid is subsequent treated of in illustrating the mode of electrifying a pages.

## Of Mitigating the Effects of Electricity.

198. The same charge of electricity, which would be duce a shock or deflagrate a wire if received through knob, may be received imperceptibly through a fine per The power of an electrical machine, or that of a be battery is thus easily paralyzed. Hence the employed of pointed rods of metal as a protection against lighter Hence, also, it is necessary, in the construction of trical apparatus, to avoid all edges or acute projected unless where it is intended to facilitate the pussage of the fluid.

199. Rationale.—When a knob is approached to use cited conductor, as soon as any portion of it is near enced to be struck, so much of the ball is at hand as to the whole discharge to take place at once; but when the apex of a pointed wire is near enough to receive a portion of the fluid, the other parts are too remote to come use action; and whether the point advances to the electrical body, or the body approximates the point, the fluid is decharged before it is within striking distance of the thicker part of the wire.

200. In fact, the difference between a discharge by a point, and through a knob, resembles that which exists be tween making an auger-hole in a reservoir to let out a

fluid, and opening a floodgate.

# Of the proper Mode of constructing and putting up lagh-

201. The competency of pointed rods to protect against lightning is dependent, not merely on the excellence of the point, which should be of platinum, but on the

de in which the joints in the conductor are made, and nature of the soil in which it terminates. If a wire, nted at one end, be blunt at the other, and the nearest iductor to the blunt end be not pointed, the charge will pass off gradually because received at a point. Moreon pounded of its own conducting power, and that of medium in which it terminates.

- 202. A metallic rod, terminating in a glass handle, in asses of sulphur or resin, or in dry sand, would not erate as a conductor. It cannot receive electricity, bease it cannot deliver it.
- 203. Moist earth is but an imperfect conductor; since owes its faculty of conducting to water, which, according to Cavendish, conducts with 200,000 times less facility an iron.
- 204. Lightning rods should, therefore, be connected, by dering, with an extensive metallic surface buried under earth; as, for instance, with sheets of lead or copper. 205. The cases in which conductors have been found competent, are, I am satisfied, referrible to their inadeate communication with the earth.
- 206. Thus is an important lesson given with respect to means of protection against lightning. In vain do we ploy for this purpose a conductor, however perfect, if it we communication with the earth only through a limited ntact with the soil, which, being at best an imperfect nductor, may become an electric by desiccation.
- 207. I have surmounted the lightning rod by which my insion is protected, by 17 copper wires pointed at one d, and at the other soldered into a hole drilled in the 1, which is constructed of iron. The juncture is surneded by a globe of zinc of about two inches in diamended by a globe of zinc of about two inches in diamended, above which the wires extend divergingly. The coper wires, by their association with zinc, are protected of oxidation; while their greater fusibility, as compared the platinum point usually employed, is compensated their number. The rod thus mounted rises about 10 st above the apex of the roof, to the copper covering of anich its lower end is soldered. The copper covering off e rain, and this pipe with those of the public water works; I the joints being made with screws or solder. By these

means a most ample communication with the earth is tained. Analogous means should be employed in them of all lightning rods, in situations where access can be to a similar ramification of metallic pipes.

# King's Improvement in the construction of Lightming Im Expedient suggested by the Author.

208. Since the preceding suggestions were made, line become acquainted with the plan of putting up light rods, pursued by King, an ingenious practical electric of Boston. Agreeably to this plan, in lieu of round is square copper rods are employed, the corners being in as jagged as possible, by cutting them with a cold chis so as to cause them to resemble rude irregular saw-to By these means it was conceived, and with reason a think, that if a discharge should be made through the greater than it could convey to the earth, the excess to be dissipated in the air, in preference to entering a in order to find a conveyance through the imperfect of the conveyance through the

ductors of which buildings are constructed.

209. This ingenious innovation has led me to com plate an expedient in which I have great confidence. would extend the rod forming the conductor, under ground, to a tree or post, at the distance of from fifty one hundred feet, and cause it there to terminate in a k of copper, of which the edge should be so cut as to rest ble large saw-teeth. These should be bent out in s manner as to project horizontally from the band, at distance of not more than six inches from the surfac the ground. It will, I think, strike any experienced e trician, that by means of such a serrated band of cop as I have described, the electric fluid would be dissipt into the air, and over the surface of the ground, will facility so great, as to prevent it from leaving the c ductor for any other channel in which it might do t chief. The portion of the conductor under ground, she consist of copper, but I think that a wire of a inch in ameter would be sufficiently large for that portion.

## Experimental Illustrations.

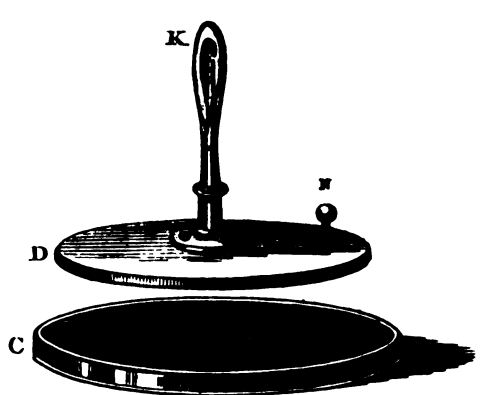
210. Effect of a point shown in drawing a charge fra battery or conductor:—proved, also, that its power

reakened or destroyed by being associated with an elecric or imperfect conductor.

ADDITIONAL MEANS OF PRODUCING ELECTRICITY.

If the Electrophorus.—Of Electricity evolved by the Friction of Caoutchouc.—Of Electricity evolved by Pressure.

Of the Electrophorus.



211. This is a peculiar instrument for the excitation of electricity, which we owe to the celebrated Volta. rical induction has already been explained (e. 107, &c.); ikewise the susceptibility of resinous substances to be elecrified negatively by friction has been illustrated. (e. 3, 33.) The electrophorus is indebted for its efficiency to the coperation of the effects of friction with the inductive wer. By rubbing the cake of resin, C, with the fur side I a dry cat skin in a dry atmosphere, the resin is negaively electrified. If the metallic disk, D, be placed upon he resin while thus excited, it will acquire a greater capazity to receive electricity, in consequence of the proximity of the negatively excited surface of the resin, as has already been explained in the instance of the condensing electrometer. (e. 127, &c.) Hence, if placed upon the resin with the aid of the glass handle, K, on applying the knuckle to the knob, N, a spark will pass, and again when

the disk is lifted about an inch or two from the resin to other spark will be received; and this may be continued in a dry air for a longer time than any operator was have patience to repeat the operation. In dry waster and in a dry apartment, an electrophorus will give part for many weeks, without being again excited by frame. It was through the aid of this property of his exception, that Volta was enabled to contrive an apparation the production of instantaneous light, of which a engraving and description will be found in the Compodium, (327, &c.)

Of Electricity evolved by the Friction of Caoutchem.

212. Dr. J. K. Mitchell has ascertained that in the seface of a bag made of caoutchouc, or that of a thin sef of the same material stretched on a frame, there is a prising susceptibility of electric excitement. The atmetive power, consequent to the excitation, appeared be peculiarly powerful.

## Of Electricity evolved by Pressure.

213. There are many substances, especially gum class or caoutchouc, which, if placed upon the disk of an determent, and pressed, will, on the cessation of the presure, evolve enough electricity to be indicated by the gold leaf.

Evolution of Electricity from Caoutchouc by Pressur-



214. C, cap of electrometer; D, but dle of metallic disk, between which and the cap of the single leaf electrometer, may be seen a piece of caoutchouc subjected to pressure. The disk being subdenly removed, the leaf will strike the ball as when otherwise electrified.

#### THEORETIC EXPLANATION OF ELECTRICAL PHENOMENA

215. It may be proper to give the rationale of the production, securition, and neutralization of electricity, agreeably to the doctrine of two feets.

216. This doctrine has, within the last thirty years, been games a credit, especially on the continent of Europe.

### Of the Theory of two Fluids.

217. According to this hypothesis, when a body is electrically excited, a neutral compound, formed of two electrical fluids, is decomposed; one exp

mtes in the exist dody, the other is evolved. That which accumulates excited glass is evolved by excited resin; and vice versa, that which tamulates in resin is evolved by glass. Hence, according to this theory, two electricities are always produced both by resin and glass; though fluids accumulated in them respectively, being, in the first instance, exactly recognised and associated with the substances in which they

coordingly.

ss cylinder or plate is rubbed against a cushion, it gives off resinous and eives vitreous electricity. This surcharge of the vitreous fluid is remed by the excited surface, until, in revolving, it comes opposite to the lector. (e. 21,28.) The change of capacity, which caused the glass to et the resinous and attract the vitreous fluid, ceasing with the friction to ich it owes its existence, an equalization of the electricities between the ss and the conductor, through the collector, ensues; by which the excitent is reduced, more or less, according to the extent of the respective rfaces. Returning to the cushion, the glass is re-excited to the same tent as at first; and, by a continuation of the process, gradually causes exchange of vitreous for resinous electricity between the conductors, til they are electrified as oppositely as the performance of the machine ill permit.

219. Under these circumstances, if a conducting communication be made tween the oppositely excited bodies, the electricities rush together, neutral-

r ensues, and a total cessation of electrical phenomena.

220. By some philosophers, the union of the two fluids is supposed to be oductive of the evolution of heat and light. The inevitable consequence at in separating, they must absorb those elements, seems not to be sufficiently contemplated; nor indeed justified by any observed refrigeration tween the surfaces of the cushion and electric, where the separation kes place. (e. 29.)

221. In charging a Leyden jar, the two coatings become, for the time ey are in communication with the conductors, respectively a part of them. hat which touches the resinous or negative conductor, therefore, gives up vitreous electricity, and receives resinous; while an opposite exchange resinous for vitreous takes place in the other coating. Meanwhile, the posite electricities are supposed to exercise a reciprocal attraction through e glass, and are thus retained upon the surfaces which they respectively cupy. Of course a removal of the coatings does not cause a removal of e charge.

222. When the positive and negative surfaces of a charged pane act multaneously on a metallic arch made to complete the circuit between em, the resinous electricity is attracted out of the end of the arc, near the rating vitreously charged, while the opposite effect takes place at the other ad. Thus, by a series of decompositions and recompositions, or by curents passing each other, the different surfaces are restored to their previous

ate.

223. The principle of induction is much resorted to, agreeably to the octrine of two fluids, to explain the passage of a spark between a body ther vitreously or resinously excited, and one which is neutral. It is suposed that the fluids attract each other in all proportions, and in a way to ring into equilibrio the self-repellent power of the homogeneous atoms ending to dissipate the aggregate fluid, and the attraction of the heterospecous atoms tending towards its accumulation. Yet it is conceived that

the equilibrium is liable to be destroyed by the presence of a best charged with either electricity. If it be vitreous, by induction a residual vitreous fluid from any mass in its vicinity, and attracts the reason is surcharge be resinous, it repels resinous electricity, and attracts the reason. When this species of reaction reaches a certain degree of intracts which, by the surcharge, and the heterogeneous electric matter, which, by the process, has been accumulated in the neighbouring conductor, run producing an electric discharge, and under favourable circumsus a spark or a shock.

224. Thus the unknown electric compound, formed by the unit is two electricities, is considered as liable to be decomposed by the premate

of either of its constituents in excess.

225. When of two portions of the same matter, one is remove, the disactually combined with a common object of attraction, I cannot missess wherefore the one should be displaced by the other.

## Additional Remarks explanatory of Franklin's Hypothems

226. According to the theory of Franklin, the effect of an electric period chine in drawing electricity from the cushion, and accumulating to prime conductor, has some analogy with that of a wheel or pump for ping water.

227. The increased capacity for the electric fluid, arising from the in the glass, operates like a bucket on a wheel, to enable it to receive the tion of fluid; and the loss of this capacity, after the friction cease, a legious to that inversion of the buckets which causes them to be empted to

every semi-revolution.

228. When the liquid thus raised is withdrawn from one reservor to thrown into another, so that the quantity in the former is the analogous to that of the cylinder or plate of an electrical machine in creating a surcharge to a the surfaces of a pane at the expense of the other. Should the return be allowed subsequently to communicate by a syphon or stone, and not sequence being on the same level, a previous equathrium of pressent restored, the other performed by the syphon would be analogous to that a metallic arc in restoring the electrical equilibrium of the charged arrays of a coated pane; excepting that the velocity of the electric current and comparably greater.

of Franklin, the messant agency of the attraction between the fluid at the glass should be kept in mand. It is necessary to suppose an interestation to exist between the glass and the electrical particles, what is tween these particles there is an intense at the pulsion. When there was are unimpeded, there can be no electric accumulation, because operant equally, they must produce an equal distribution of the electric fluid.

230. In a charged pane, this tendency to an equilibrium is repersive by the impermeability of the glass, so that when by the operate of the machine, the fluid is abstracted from one sale of the pane to be piled upon the other, the same attraction of the glass which resists the withdraws of the fluid from the negative surface, retains it upon the positive surface soon as it reaches it; while the surcharge thus retained, prevents any accession to the negative surface of electricity from other bodies. Thus an electrical surcharge in a coated jar or pane, is dependent on its attraction for the

gative surface, and the repulsion between it and the electrical matter in ser bodies.

231. In concluding this theoretic explanation, I refer the reader to some ggestions in my letter to Faraday, which are quoted at the close of my reatise on Electro-magnetism, page 77.

## Rationale of Electric Light and Ignition.

232. It appears evident that there is a re-action between

eat, light, and electricity.

- 233. It is not surprising, therefore, that the presence of ne should cause the appearance of either of the others; nce they all, evidently, pervade nature. It may be concerned that the electric fluid is luminous when projected ith intensity into the air, in consequence of its carrying long with it the light encountered in its progress through onderable matter. In like manner it may cause the excication of caloric, by displacing it when latent; or by dding temporarily to its repellent power, it may enable to overcome the attraction of cohesion; in which case metal, no doubt, contains caloric enough to produce a iolent, or even explosive separation of the metallic parcles. (Compendium, 274, 275.)
- 234. It has been mentioned that, agreeably to the late researches of Faday, the electrical spark varies both as to its dimensions and shape, with
- e gas employed as the medium for the experiment. 235. "In nitrogen, it is alleged to have been found pre-eminently beauful, having still more of the purple hue than in air." "In oxygen the parks were whiter than in air, or nitrogen, but less brilliant." "In hydroen they had a fine crimson colour not due to its rarity," since rarefaction iminished this characteristic. "In carbonic acid the spark was similar to rat in air, but had a little green in it. The sparks were remarkably irreular in form, more so than in common air, and could be obtained longer, nder the same circumstances, as to size of ball." "In muriatic acid the park was bright throughout, and nearly white, never presenting any dark arts as in air and some other gases." "In coal gas the spark was somemes green, sometimes red, and occasionally one part was green and anoher red." "These varieties of character," says the distinguished author, impress my mind with a feeling that they are due to a direct relation of he dialectric, through which the discharge occurs, and are not mere results of the casual ignition or secondary kind of action of the electricity upon the marticles which it finds in its course."
- 236. From the facts cited above, it must be evident that a specific influence is exerted, by gaseous substances, upon the form, dimensions, and coour of the electric spark; and from some paragraphs which I subjoin, this appears to be equally true as respects the brush; yet when the diversity of colour produced by different substances, by reflection or transmission; also from their combustion, or from their mere presence, as in the case of stron-

tia (1493), when no chemical change is experienced or induced, a feet appear to me that the variations in the hue of the light conined, plausibility of the rationale given above by me.

237. Air.—Fine positive brushes are easily obtained in air at common runal possess the well-known purplish light. When the air is rarefied to min tions are very long, filling the globe, the light is greatly increased, and a situal

tiful purple colour, with an occasional rose tint in it.

238. Origen.—At common pressures, the brush is very close and compress of a dull whitish colour. In rarefied oxygen, the form and appearance in the colour somewhat purplish, but all the characters very poor compared to an

239. Nitrogen gives brushes with great facility at the positive surface, is well any other gas I have tried they are almost always fine in form, light, and six and in rarefied nitrogen are magnificent. They surpass the discharges in an extension of the charges in the charges in an extension of the charges in 
gas as to the quantity of light evolved.

240 Hydrogen, at common pressures, gave a better brush than cryrea was not equal introgen; the colour was greenish gray. In rarefied hydroges in fications were very fine in form and distinctness, but pale in colour, with a case velvety appearance, and not at all equal to those in nitrogen. In the rares and

the gas, the colour of the light was a pale gray green.

241 Coal gas.—The brushes were rather difficult to produce, the contract mitrogen being great in this respect. They were short and strong, geom. 41 greenish colour, and possessing much of the spark character for, occur at a left the positive and negative terminations, often when there was a dark interest of length between the two brushes, still the quick, sharp sound of the spark tage. duced, as if the discharge had been sudden through this gas, and partaking the respect, of the character of a spark. In rare coal gas, the break forms were too.

but the light very poor and the colour gray.

242. Carbonic acid gas produces a very poor brush at common pressure.

gards either size, light, or colour; and this is probably connected with the brain. which this gas has to discharge the electricity as a spark. In rarefied carries at the brush is better in form, but weak as to light, being of a duil greened as purch hue, varying with the pressure and other circumstances

243. Mariatic acid gas —It is very difficult to obtain the brush in the gas it ex mon pressures. On gradually increasing the distance of the rounded was sparks suddenly ceased when the interval was about an inch, and the interpretability. which was still through the gas in the globe, was silent and dark very short brush could for a few moments be obtained, but it quickly discussed Even when the intermitting spark current from the machine was used of the only with difficulty obtain a brush, and that very short, though I used me rounded terminations (about 0.25 of an inch in diameter) which had be the them most freely in air and bitrogen. During the time of this difference is muriatic gas, magnificent brushes were passing off from different parts of the chine into the surrounding air. On rarefying the gas, the formation of the was was facilitated, but it was generally of a low squart form, very poor in light and minutes on both the positive and negative surfaces. On rarefying the gas states a few large ramifications were obtained of a pale bluish colour, utterly unless the in nitrogen.

MEANS OF ELECTRIFYING PATIENTS EITHER WITH SPARES CO.
BY SHOCKS

## Electrifying with Sparks.

244. A person, seated on an insulated chair, is made to communicate with one of the conductors. Being thus acgatively or positively electrified, sparks may be taken from any part of the body, by a metallic knob or point. If the sparks emitted by the knob be too severe, the point may be used; and if the current from this be too powerful, it

r be covered by a wooden cone, C, represented in the twing engraving.



Apparatus for Electrifying by Shocks.



245. In order to subject a person to shocks, a coated is used with two knobs; one in communication with the ide coating through the rod, R, which supports it, the er supported on an insulated wire, W, so that it may made to approach or recede from the knob which comnicates with the inner coating. To the outer coating I the insulated knob, chains are attached, each termining at one end in a knob of metal with an insulating

handle. The handles are held by the operator, and knobs applied to the patient, so as to have between the part to be electrified. The coatings of the jar has everally connected with the different conductors and electrical machine, the charge increases in the jar makes become strong enough to strike through the interest tween the knob connected with the innner coating that insulated in its vicinity. Of course, the height of charge depends upon the interval thus left, according the operator's discretion, or the feelings and fortital the patient.

## OF GALVANISM,

OR

## VOLTAIC ELECTRICITY.

ajority of electricians, Dufay's suggestion of the existence two fluids is sanctioned (E, 217, &c.\*), while, by others, makin's idea of one fluid is preferred; yet philosophers high pretensions question the existence of any imponable matter. Whewell, of the University of Cambridge, England, published an essay a few years since, of which was the object to prove, that "all matter is heavy." raday, so highly distinguished by his Herculean resches, advanced the idea that electrical induction is the sequence of an action of contiguous ponderable partision an affection arising from their association with onderable matter. In letters published in Silliman's rnal, I consider myself as having invalidated the positive taken by these eminent philosophers.

that there is any transfer of imponderable matter ing electrical discharges. Latterly I have conjectured, electrical phenomena are due to the opposite polarition of electric matter, which pervades all bodies and all electric matter in different states of polarization, and an electric matter in different states of polarization, and an electric current consists of a succession of polarizing impulses. It is that the two electric matter in different states of polarization impulses. It is that the two electric matter in different states of polarization impulses. It is that the two electric matter in different states of polarization in the matter in different states of polarization in the matter in different states of polarization.

ference is founded shall be the su 302. Nevertheless, it seems ur pedient to suppose, provisionally lectric imponderable matter; and ne current are more easy to con

ersally to be deemed existence of fluid he movements of of and describe,

e on "Electricity Prolium, the letter C.

In referring to the numbers of paragraphs in the er," I shall prefix the letter E: in referring to my

than those of two fluids, the language founded of Franklinian theory is much used, even by those who

sume the existence of two fluids.

there must, in every part of any active electric curse, two fluids flowing oppositely to each other. The quest of either fluid must be increased as that of the others minished, so that there can neither be any diminuted any increase of the sum of the quantities of both is Moreover, the current in an electric circuit can be be directed from positive to negative, than from negative, or minus and plus, are applied to the states of bod implying, of course, a deficiency in the one case, and a dundancy in the other. Moreover, arrows indicating current to move from the positive to the negative pole almost universally used.

responding with the theory of Franklin, wishing it to understood, that when employing the words electrical rent, as designating the phenomena of an active galvereut, I do not mean to convey the idea that those ware sufficiently descriptive or accurate. The words are sufficiently descriptive or accurate. The words are sufficiently descriptive or accurate to designate unknown cause of certain effects, which are in some spects such as might result from a current of imponder

matter.

Of the Elementary Battery, or Simple Galvanic Pair. its Circuit: more properly designated as a Galvariad.

305. It is generally known that metals are liable what is called corrosion, arising from a reciprocal act or reaction, with certain gases or liquids; also, that s metals will react with liquids or gases, with which o

metals will undergo no reaction.

306. It may be assumed as a general law, that we two metals are simultaneously exposed to a liquid, excising with one of them a more energetic reaction to with the other, the phenomena will be such as to come with the idea of an electrical current from the more ceptible metal, through the liquid, to the other metal:

rom this, through any competent conductor, back again o the first mentioned metal.\* To this result a complete :ircuit is indispensable; and it is on the completion of the sircuit that the phenomena take place which are assumed o demonstrate a current. It is on this account that when he tongue is situated between a plate of copper and a date of zinc, no sensation is perceived until the projecting extremity of one, touches that of the other: but as soon as his contact is effected, a discharge takes place from the inc to the copper (or silver) disc, which affects the taste Where the surfaces are omewhat like a feeble acid. arge, the current may be indicated by the ignition of a vire made the medium of communication, or by the atraction and polarization of iron filings; but among the nost competent means of detecting such currents, is the alvanometer or multiplier of Schweiger. In this a magetic needle, situated within a coil of wire as represented

in figure 1, by its deflection detects the existence of the most feeble galvanic current, which may be made to pass through the coil by means of a

ue communication with the ends, a b, of the wire.

307. Hereafter the reaction of the galvanoelectric current, so called, with the cause of
magnetism, will be more fully brought into
view, while at present it may be well to advert to another galvanoscope, which owes its
existence to the same source. Fig. 2 represents this galvanoscope. When a small ribbon
of gold leaf situated between the legs, N S, of
the magnet, D, and parallel thereto, is made the
medium of a minute galvanic discharge, the
ribbon is swayed one way or the other, according to the direction of the current.

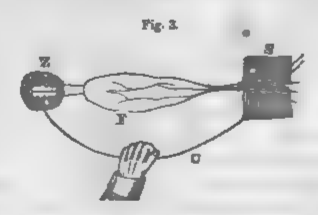
308. The tongue has been mentioned above as performing the office of a galvanoscope. Originally, the limb of a frog, with the lumbar

or spinal nerves denuded, was employed by Galvani and his nephew, Aldini; and, recently, this organic instrument has been found by Matteucci, a distinguished modern in-

<sup>&</sup>quot; I here state the opinion generally held, yet Sturgeon alleges, that in the case of iron and amalgamated zinc, although the former is much the most rapidly dissolved, it nevertheless acts as copper.

vestigator, to be competent to the detection of a const too feeble to affect a most delicate galvanometer, of a construction analogous to that above illustrated.

309. The following figure will give an idea of a far prepared so as to answer as a galvanoscope. The nuded lumbar nerves, with a portion of the spine attacks.



are represented at z, the feet at s. Now, if z be a plan of zinc, and s a plate of silver, copper, or platina making or breaking the circuit by the metallic arch convulsions will be produced. When the interruption and renewals of contact at one of the extremities of the arch are rapid, the agitation is incessant.

310. When the frog is used as a galvanoscope, the attallic plates should be perfectly homogeneous, so as a themselves to produce no discharge. In lieu of resting a metallic plates, the nerves and feet may each be enveloped

in tin, or such lead foil as is used in tea chests.

311. Thus prepared, a dissected frog may be included in any circuit, in lieu of the galvanoscopes above described. Under those circumstances, the passage of the most menute galvano-electric current will be indicated by a corresponding tremulous motion in the organic instrument employed.

312. When a galvano-electric current is produced by two metallic plates, or any other competent masses, and an interposed exciting agent, they are usually called a galvanic pair: but as three elements are requisite to the characteristic effect, I prefer to call the apparatus which they

form, a galvanic triad (306).

313. In the construction of galvanic triads, various forms may be given to the metals upon which the house is made to operate; and any two metals will be productive of analogous results, provided that one be more succeptible of erosion by the exciting liquid than the other-

preëminent degree, effice y with cheapness. Yet siler would be preferable to copper, gold to silver, and pla-

inum probably to gold.

ith respect to copper and zinc, will apply, more or less, a the case of other metals. Hence C and Z, the initials of copper and zinc, may be considered severally as resenting two metals, one of greater, the other of inferior susceptibility of erosion. The electric relation of these netals may be changed by varying the exciting liquid. Thus, for instance, in a galvanic arrangement, in which copper and zinc should be subjected to ammonia, which acts upon copper more than upon zinc, the latter metal would, in a simple galvanic circuit, give electricity to the former. It appears, also, that in lieu of two metals and one liquid, we may employ two liquids with one metal; or one metal with one liquid may be employed, provided the surfaces be in a different state as to roughness, since a more powerful reaction takes place with the rougher surface.

315. The energy of the current varies also with the greater or less disposition of the negative, or less oxidizable metal, to become tarnished by oxidation, or depositions from the liquid, and of being partially dissolved and precipitated on the other metal. These evils are so great in the case of zinc and copper, as to cause a battery constructed of them, without any remedial contrivance, to lose almost all its efficiency, before the zinc is exhausted. The zinc becomes so completely coated by a mixture\* of its suboxide with metallic copper, that due access of the acid is prevented.

316. As either silver, gold, or platinum, differ from zinc more than copper does, as to liability to erosion, were it not for their superior costliness they would be preferable to this last mentioned metal. Platinum is preëminently preferable, being less susceptible of change than gold. A minute quantity of this metal, precipitated on the surface of silver or copper, has been found to make these metals

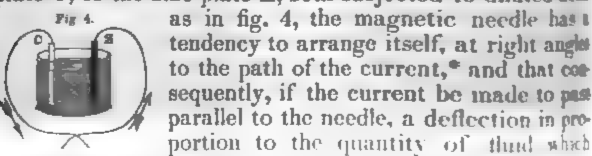
This mixture, after being softened by soaking in water, and then scraped off, I found, in one instead, to grow hot spontaneously. Probably this arose from a galvano-chemical result in between the moisture and the copper and zinc particles, during which thes were acquiring an additional quantity of oxygen.

more efficacious in a galvanic triad, by diminishing the injurious tendency to coherence, between them and hypergen. Recently it has been found, that either plumber coke, or the indurated carbon sublimed in verses we evolving gas from bituminous coal, may be substituted in the copper in a triad.

317. From the phenomena of electrical fishes, those observed by Matteucci in the muscles of this fishes, and warm blooded animals, it is evident that the law, which has been assumed to be in operation in the case of metals, must prevail no less where organic as

stances are solely concerned (306).

318. Although at this stage of our illustration, any tempt to explain the theory or construction of the metaphier, or galvanometer, would be premature, it will the highly advantageous to employ it as the best means of the tecting the direction and force of a galvanic current. It present, it will be sufficient to state, that when a galvant discharge takes place through a wire, as from the copposite C, to the zinc plate Z, both subjected to diluted and



passes, will ensue. The interposition of the coil of the galvanometer, merely reiterates the influence of the current, by causing it to circulate many times about the needle. See Electro-magnetism, for various engravings and

descriptions of this instrument.

319. The direction of the galvanic current is known from the manner in which the needle is deflected; such when the current is above, and flows in a direction parallel to the meridian, from south to north, the north pole is deflected so as to point westward, and of course the south pole, oppositely deflected, points eastward. When a similar current is flowing in the same direction below the needle, the deflection is the reverse of that just described.

<sup>\*</sup> More accurately, it may be said to take the attitude of a tangent to the carreference of a cylinder, in the axis of which the current flows.

part in any case, reversing the direction of the current, re-

320. In an elementary battery, or triad, consisting of copper, zinc and water, or any aqueous solution, the oxilation of the zinc is, as already suggested, indispensable the production of the phenomena, which are conventionally attributed to an electrical current circulating from the zinc, through the liquid to the copper, and back again to the zinc. Under these circumstances, as the oxygen of the water combines with the zinc, it might be expected, that hydrogen, the other elementary ingredient of the liquid, would be liberated at the surface of the metal, as is known to be the fact, when zinc in an impure state, as found in commerce, is subjected to diluted sulphuric acid. But, in fact, so long as there is a metallic communication between the plates, the hydrogen will be liberated only at the surface of the copper. According to Faraday, there is no reaction and consequent evolution of hydrogen, when zinc, purified by distillation, is subjected, per se, to diluted sulphuric acid; nor from commercial zinc, if the surface be amalgamated, or in other words, coated by a film of mercury. The effect of this coating is to prevent the formation of local circuits, between the zinc and certain metallic impurities which it contains. A homogeneous substance is produced, which, per se, is scarcely susceptible of reaction with diluted sulphuric acid.

321. But although neither pure zinc, nor impure zinc when amalgamated, has, per se, any reaction with the water of diluted sulphuric acid, yet if touched by a piece of copper previously immersed in the solution, the oxygen of the water combines more or less with the zinc, while the hydrogen is copiously evolved from the cupreous sur-

face.

322. This may be considered as another of the characteristic features of a galvanic circuit, though it cannot be so conveniently nor satisfactorily made manifest as the indications above cited (306, 307, 309).

323. The characteristic phenomena which have been described as resulting from a galvanic triad, are not inseparately associated with any particular forms of the generating surfaces, or arrangement of them. The same extent of superficies may be more efficacious when arranged in one way than in another, or may vary in its efficacy

with the energy of the exciting liquid; but whether citar or both of the metals be used in one plate, or in seven does not alter the characteristic features of the circus se they have been described.

#### Of the Calorimotor.

324. In 1818 I contrived a modification of the galvanic triad or tary battery, which I designated as a calorimotor or heat mover, unit to idea that a current of heat no less than one of electricity, is an effect of

galvano motive power.\*

325. The calorimotor consisted of twenty-one plates of copper consisted and twenty plates of zinc similarly exactly the plates of the one set being alternated between those of the other. Is apparatus thus constructed, the zinc is every where opposed by coperate that both of its surfaces become equally efficacious. Still this expense although consisting of forty-one metallic sheets, and twenty intersers plete with the exciting liquid, comprises only the three elements of a plete.

The ignition and deflagration of wires by the galvanic current has by phine phers generally, been ascribed solely to electricity; the heat and light result having been treated as secondary effects. But as, agreeably to the observated Davy, sanctioned by general experience, a finite portion of wire, expected in the we take circuit, may be kept ignited for an unlimited time, it follows, that if heat is me terial, it cannot be supplied by the wire, and must be supposed to flow from the monotone as the accompanying electricity. Considering the imponderable fluid into circulation by galvanic reaction, as consisting of caloric as well as electricity inferred that the proportion of the caloric increased with the size of the part the of the electricity with the number, and that in a large apparatus of one per the latter fluid was in a minimum proportion, so that it was mainly caloric, which we made to circulate by a large galvanic pair which I had constructed

latter fluid was in a minimum proportion, so that it was mainly caloric, which we made to circulate by a large galvanic pair which I had constructed.

It is demonstrated by the phenomena of thermo-electricity, of which as accordingly will be given in due course, that a current of heat may be the primity rune to galvanic current, in thermo electric apparatus, just as much as a correct of recity is the cause of an evolution of heat from an apparatus, such as above due to called hydro-electric, because a liquid is requisite to the result. It seems there in reasonable to consider, that both caloric and electricity may be products either a galvane or thermo-electric reaction, and that if heat he a secondary product a consider.

case, electricity is no less so in the other. There is the most ample proof that the cause of the one, as well as of the other, exists in metals to an enormous der to me, increase of electricity, are, it seems to me, more difficult to answer than any that can be adduced against the existent of a material cause of cal rific repulsion.

Contemplating a galvanic pair as simply a mover of the electric fluid. Volta had called such pairs, as existing in his series, electromotors. By analogy, I applied the

term calorimotor to signify the large pair or heat mover above mentioned

The opinion above alinded to, that the heat of an active galvanic circuit is not a electrical effect, as generally supposed, but that on the contrary, caloric and electrical effect, as generally supposed, but that on the contrary, caloric and electrical products of galvanic reactions was entertained when the causes, both of the calorific and electrical phenomena, were generally considered as material fluids, the electro magnetic powers of the circuit in question being the known. It will, in due course, be explained, that I am now disposed to considered powers of the active circuit as the consequence of waves of polarization, and never bly of concomitant vibrations or inovenients, in one or more imponderable principles, which, pervading all space, are combined more or less in a condensed state will ponderable atoms, and enter enormously into the constitution of perfect conditions in other words, metals.

But this view of the subject is entirely cans stent with the inference, that had and electricity are connate and collateral products of galvanic reaction, and the heat is no more an effect of electricity, than electricity is an effect of heat.

vanic triad, one associated zinc surface, one associated copper surface, and

he liquid interposed every where between them.

826. I ought to mention, that by means of a silver thimble, within which is minute piece of zinc was supported and insulated, Wollaston made an dementary battery capable of igniting a minute platina wire. In enlarging he size of the galvanic triad, as above stated, my course was diametrically pposite. Several methods of accomplishing this design occurred; among thers, I contemplated concentric coils of copper and zinc, the former being wound as to surround the zinc completely. Also, to unite several sheets opper to one bar of metal, and several zinc sheets to another bar, the sinc and copper alternating. To the latter plan I gave the preference. I wenty sheets of copper, and a like number of zinc sheets, associated as above described, were secured in a wooden frame, so as to leave about a quarter of an inch between them. One end of a rope, passing over a pulsy, was attached to the frame, while to the other end a counter weight was astened. A cubical box of suitable size being placed under the frame thus suspended, and supplied with diluted sulphuric acid, the frame could be owered into the acid liquid, or lifted out in an instant.

827. When one plate of zinc is placed between two of copper, as in the Wollaston elementary battery, represented by the adjoining cut, out of four

copper surfaces two are useless; but in a calorimotor where the same quantity of zinc is used so as to form several plates, say four, for instance, five copper plates of the same size, will be sufficient to constitute a calorimotor in which each zinc plate will be opposed by a cupreous surface, and only the outermost surfaces, the two external copper plates, will be useless. Hence, of the ten surfaces in the five copper plates, all but two are brought into action.

328. A battery, resembling the calorimotor in the alternation of surfaces, has lately been constructed, in which iron plates are made to perform the office of those of copper: much merit is attached to the construction, no

reference being made to its previous employment by me.

329. While I was engaged in these experiments, Mr. Lukens, to whom I had mentioned the various plans which I had projected, of calorimotors, put into operation, with the aid of Dr. Patterson, the plan of concentric coils. Although this apparatus had less surface than that which I had constructed, and would not heat as large a wire as the latter, yet it communicated a much higher ignition to a small wire than mine was capable of producing.

330. Soon after this experiment of Patterson and Lukens, J. P. Wetherill, Esq., had a pair made in the form of concentric coils, of one hundred square feet of surface. From this great results were expected, and a large assemblage, of those interested in such experiments, attended to see the mammoth pair tried. But to the surprise of all present, it did not induce even a red heat in connecting wires, such as had been vividly ignited by my apparent.

ratus, or that of Patterson and Lukens.\*

Bome time after these large pairs were constructed, similar apparatus was made by Mr. Pepys, of London, and likewise by Col. Offerhaus, on the continent; and although a memoir, in which I gave an account of these experiments, was published in Silliman's Journal, and in the Annals of Philosophy, Pepys' apparatus of concentric coils has been treated of as an original contrivance in Turner's Chemistry, and elsewhere. The apparatus of Offerhaus, though made long after those constructed

831. It struck me afterwards, that in a very large pair, the intensity not increase with the quantity of the fluid generated, since the imposite matter, or the waves of polarization, cannot be concentrated a given small point of afflux, as soon from a large surface, as from the one.

832. It is analogous to the case of a small and a large stratum of t

by Petterson and Lukens, and by Wetherill, was described as if it had bout

form of apparatus.

Finding that in consequence of the low intensity of the current from my motor, consisting of one pair formed of two surfaces of fifty equare feet, it is produce a heat sufficient for the active deflagration of wires, I subsequent verted it into an apparatus of two pairs, the copper sheets of one alterests the zinc sheets of the other

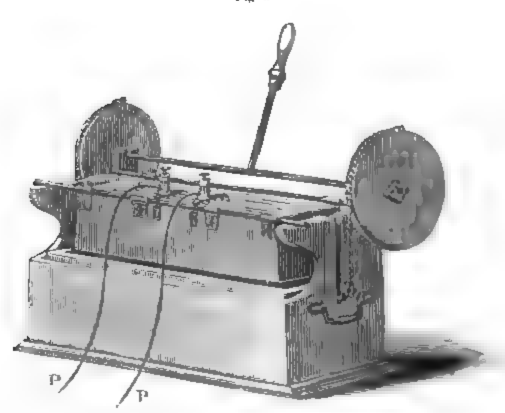
Two of the surfaces were united by a mass of solder. To the others forces soldered, into which the ends of the wire were fastened, which was to be medium of discharge. This form of the calorimeter I have found of great we means of igniting gaseous mixtures in outliometrical experiments, and other

cosees requiring ignition in close vessels

Figure 5 represents a calorimotor of four pairs, so constructed that the dissectation in a vessel of suitable form and size is brought into contact with tallic surfaces, by applying the hand duly to the end of a lever associated with pulleys and two cords, attached severally to the ends of the vessel. The experience wound up on the pulleys, raise the copper vessel so as to cause the pair immersed in the diluted acid.

The two forms of the calorimotor represented by fig. G and 7, have less used by me for what is described in my Compendium as "galeano ignition." Within any cavity, ignition of any intensity short of fusing platina may be possibly making a platina wire the subject of a galvanic discharge from an instruthis kind. I first resorted to this process in the year 1820, for the purpose of the gaseous mixtures in sudiometers of various forms. In June, 1831, I applied nite gunpowder in rock blasting; and to this object it was subsequently agreeably to my recommendation, by Colonel Pauley, Professor O'Shornes others.

Engraving and Description of a Calvrimotor of Four Pairs, or Truds
Fig. 6



noth of one depth. It is well known that the surface in both, being sustained at the same height above the points of efflux, a jet from the former naratum will go as far as one of the same diameter, emitted by the latter.

21 383. There was, however, one characteristic noticed by me, in which his analogy does not hold; I allude to the fact, that fine wire will not fuse when made the medium of discharge from a very large pair, although the

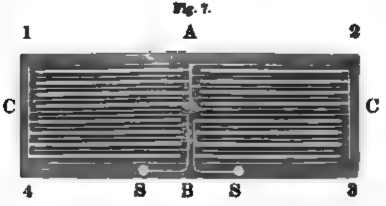
name wire may be fused by a small one.

2011. The capillary wire ignited by the thimble battery of Wollaston, 2011 not be similarly ignited by a calorimotor of fifty square feet. I have an amount one of that the coil of Patterson and Lukens ignited an iron wire about No. 24, more intensely than my larger apparatus of one pair, consisting of

"**alternate** plates.

885. Adopting the idea, that a current of electricity consists of waves of polarization, the mystery may be thus explained. The application of a wire to the electrodes, disproportionate in sectional area to the triad, the sircuit of which it is employed to close, impedes the polarization of the greater part of the masses which enter into the construction of the triad. Hence the polarizable matter within those masses is prevented from attaining even that low degree of tension, of which one triad is susceptible when its circuit is completed by a competent conductor. It appears, therefore, reasonable to suppose, that the waves in a large calorimotor move less





This machine consists of sixteen plates of sine, and twenty plates of copper, each twelve inches by seven, arranged in four galvanic pairs. The plates are supported within a box with a central partition of wood, A B, dividing it into two compartments. Each of these may be considered as separated into two subdivisions, by four plates of copper between the letters C C. Of course the box may be considered as comprising four distinct spaces, No. 1, No. 2, No. 3, and No. 4. The circuit is established in the following manner. Between the sine plates of compartment No. 1, and the copper plates of compartment No. 2, a metallic communication is produced, by soldering their neighbouring corners to a common mass of solder, with which a groove in the wooden partition between them is filled. With similar masses of solder, two grooves severally made in the upper edges of each end of the box are supplied. To one of them, the corners of all the copper plates of space No. 1, and the sine of space No. 4, are soldered. To the other, the sine plates of space No. 2, and the copper plates of apace No. 3, are soldered in like manner. Lastly, the sine plates of No. 3 are connected by solder in a groove, and the copper plates of No. 4 are in like manner connected by solder in another groove. Upon the ends, & S, of the solder just mentioned, the gallows screws are severally soldered, and to these the rods, P P, called poles, are fastened. The means by which the acid is made to act upon the plates, must be sufficiently evident from inspection. Depressing the handle causes the wheels to revolve, and thus, by means of the cord which works in their grooved circumferences, to lift the receptacle which holds the acid, until this occupies the interstices between the plates.

The rationale of the increase of intensity resulting from the above described con-

struction, will be given under the head of compound circults.

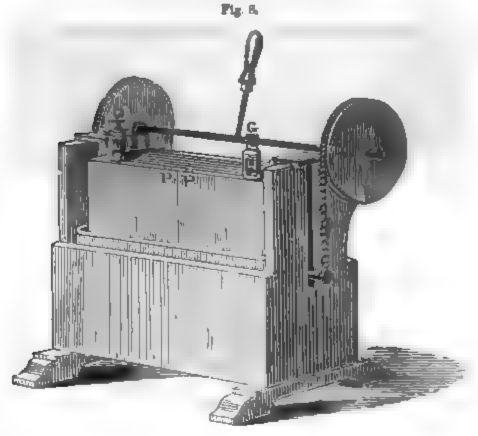
speedily, while they have, on the average, much further to go that the created in comparatively minute apparatus, essentially of the mass struction.

336. There seems to be an unalogy between the part performed by conductor in completing a galvanic circuit, and that of a kerrer repleting the circuit of a magnet so far as this; that in either case, to in by which the completion is accomplished, is the cause as wall as the ject of the resulting power.

#### Of the Galvanoscopic Frog.

337. It must follow, from the facts already alleded a that the limb of a frog must combine all that is essential to a galvanic triad; and, agreeably to the observations Matteucci,\* sections of the muscles either of warm cold-blooded animals are competent to give discharge analogous to those of a very feeble galvanic triad.





The apparatus represented by the preceding figure is a calorimotor quite and 💝 in construction to that above described, but which differs from it in be agone-fourth of the size, and in consisting of two pairs instead of four. The pairs in nine inches by seven. There are four of zinc, and six of copper.

In the form of sheets arranged as two pairs alternating, Faraday, in some of the second s

researches, employed a calorimotor

Of course, when consisting of more than one triad, the calorimeter ceases to a under the head of simple circuits. It operates, when constituted of mire than " triad, upon the principle of a compound circuit, which will be explained in the terchapter.

Annales de Chymie et de Physique, Trosseme Serie, Tom. 7, page 455.

shocks to produce ignition, or deviations of the galvanometrical needle, are, no doubt, a series of galvanic triads, in which organic masses, associated in a series voltaically, perform the office of the plates in galvanic triads, forming a compound circuit, of which the description and explanation is the subject of the next chapter.

## Experimental Illustration.

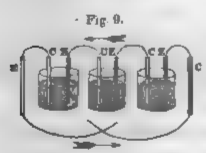
- 338. The electrical excitement arising from the contact of copper and zinc, shown by means of the single leaf electrometer.
- 339. Effect of the active circuit upon Schweiger's galvanometer and the gold leaf galvanoscope. Also, upon a leech or prepared frog.
- 340. Disks of zinc, provided for the purpose of producing a galvanic discharge through the tongue, when severally placed over and under it; the projecting limb of each disk being brought into contact.
- 341. A plate of copper and a plate of zinc, connected by a small wire, being plunged into a vessel containing a lilute acid, the ignition of the wire is produced. The same result obtained, by concentric coils of sheet copper and sheet zinc: also by several plates of copper and zinc, alternated with each other; all the copper plates being associated by one metallic bar, and all the zinc plates by another. These heterogeneous surfaces have, every where, nterstices between them, and are connected only by a small wire, which is deflagrated as soon as they are impersed in an acid.

## CHAPTER II.

OF COMPOUND CIRCUITS, OR VOLTAIC SERIES.

342. Having briefly described the construction and the process of a single galvanic triad, or elementary battery as it is usually called, I will, in the next place, direct at-

two or more triads. For the illustration of this and complicated form of the galvanic battery, a series of the triads will answer every purpose, since, were any make however great, interposed between the terminating transported would be in a situation, theoretically and practically analogous to the triad occupying the mean an in a triple series.



343. Let us suppose three galanticade like that represented by Fig. to be situated side by side, no metal communication existing between plates of any one triad, while a plate in each communicates, by at tallic conductor, with one of a differ

metal in one of the triads on either side; also, let the treme plates communicate by wires, meeting as resented.

344. The plates of neither triad communicating dire neither can form an independent circuit, but still an them may make a discharge through the complex con tor, formed by the uniting wires, and the intervening tions of the exciting liquid. It follows, that if the plates in the exterior receptacles were both of one me so as to be incapable of producing a galvanic current. middle triad would act as a simple elementary batter. forming its circuit through a complicated conductor. as in the actual construction, the plates of each triadar milarly excited, the disposition to discharge electricity ists in each. In other words, each copper plate besurcharged, each zine plate deficient, to a communiextent.\* Hence, the redundancy engendered in the c per plates, is neutralized by the deficiency engender the zinc plates; and no more of the fluid can pass at point which may be selected, than that which is gener by the two plates, between which the selected point or Intensity is all that is gained in the complex circuit. all the pairs have to use one complicated errout, they cooperate to effect a current in that circuit.

It should be recollected that the zine, discharging through the high the per, renders this metal proportionably redundant, and becomes proportional tenent

charge from the zinc to the copper, through the liquid, is compensated by that of the copper to the zinc, through the wire, excepting so far as the current is retarded by the robstruction created by the nature or dimensions of the

"conductor interposed.

346. In the compound circuit, the discharge from the zinc to the copper of each triad, through the liquid, is compensated by the discharge from the copper of one pair to the zinc of the other, and as the deficiency in each zinc plate is exactly equivalent to the surcharge produced in each copper plate, it is evident that the equilibrium must be restored almost as soon as destroyed, where the conducting communication between each pair and that between the extremities is very ample. But after making the communications ample every where else, if there be a deficiency arising either from the size, or the nature of the interposed body, between any two of the triads, the whole of the series will cooperate to effect a discharge through it. So long as this cannot be accomplished, the copper and zinc plate of every triad is brought into an opposite state of polarity, producing a tension, or in other words, an effort to effect a discharge. Hence, the intensity of the effort to overcome the obstruction, is as the number of triads, excepting the loss by retroconduction. This loss necessarily augments in an increasing ratio, so that there is probably a point, at which any numeric extension would injure by promoting retroconduction, more than it would contribute to the intensity.

347. Usually, a galvanic series is considered as composed merely of pairs, each consisting of an electro-negative and an electro-positive metal, copper and zinc, for instance. This was the idea of the inventor, who erroneously ascribed the energy of his apparatus to an electro-motive power in the metals, treating the liquid interposed as performing only the humble office of transmitting the electricity from one of the electro-motive surfaces to the

other.

348. The first construction of the Voltaic series was that of a pile of such pairs, separated from each other by moistened cloth or pasteboard. Hence, the appellation Voltaic pile is often used to signify a series of galvanic triads in any form. Fig. 10 may give an idea of the cele-

774.10

brated pile of Volta, which has immortalized in name of the inventor, and given rise to rest unsurpassed in importance by those of any other invention. The first four, and the last two use and copper plates, are distinguished by the letters, C Z. By these means the relative position of all the plates may be understood, as we set those of the disks of cloth represented by the vening dark lines.

letter to the president of the Royal Philosophical Society of Great Brown 350. "The principal result is the construction of an apparatus which is respects the shocks it is capable of giving to the arm, resembles that of a Leyden jar, or of an electrical battery feebly charged, but which seel # incessantly, being self charged after each explosion; which is, in isc. dowed with an inexhaustible charge, or perpetual action or impulsion, and the electric fluid, but which is peculiar in this power of continuous acts. and, moreover, is peculiar in not consisting, like the ordinary batteres of one or more jars coated by conductors. The apparatus of which I was, an assemblage of good conductors of different kinds, arranged in a crisis manner. Twenty, forty, or sixty, pieces of copper (or better of sour) each applied to a piece of tin (or much better of zinc), and an equal seber of strata of water, brine, ley (or other solution which coolings and thun water alone), or pieces of card, skin, &c., sonked with any of the liquids; these being interposed between each couple of the different metals. alternately, so as to have the three kinds of conductors arranged is the same order throughout. Such is the constitution of the new instruction This instrument imitates, as I have alleged, the effect of Levden und batteries, in giving similar shocks, which are, nevertheless, is cores either by the noise or force of the explosion, the length of the spire A very inferior in intensity to those given by batteries of coard and a second

349. The following is the account given of his invention by Volta, 41

351. "I propose to call this instrument the artificial electrical organ, as having both in principle and form, as I have constructed it, much grant resemblance to the electrical organ of the torpedo, than any other known electrical apparatus. In truth, it is no less than the latter, composed these

highly charged. Although equivalent only to a battery of am new of face, very feebly excited, the pile has the precionent virtue of is the coto to be charged in advance, by an electrical machine, in order to give these

ly of conductors."

as often as it may be properly touched.

352. After giving minute instructions for the construction of his pile. \*\*
author states, "should it comprise twenty of the metallic pairs, it was a
only be competent to cause the leaves of a condensing electrometer? 4verge, but will give a charge adequate to a spark, and to create a \*\* 552' of
in the fingers, as often as they may be brought into contact with the creamittes simultaneously, resembling that of a torpedo, which has been of
tremely enfeebled."

353. It is remarkable, that Volta in this communication took not be smallest notice of the rapid decline of power, which ensues in his apparatus and the fact that it soon becomes comparatively effect. The only recognized

that being caused by the evaporation of the moisture, it might be prevented by a resinous covering. But the inadequacy of the escape of moisture to axplain the diminution of power, ought to have been sufficiently evident from the striking fact, that this diminution ensues in the couronne destances. In this apparatus, as well as in the pile, the acid becomes saturated, and the plates coated with reduced copper and suboxide of zinc, although, from the ample supply of water, there can be no injurious desiccation.

#### Of the True Members of a Voltaic Series.

354. I have endeavoured to make it evident, that the apparatus, which has been called an elementary battery, or galvanic pair, is more properly considered and designated as a galvanic triad; since to form it requires, at least, three agents. Consistently, the metallic pair, belonging to each triad, comprised in a voltaic series, does not consist of the plates which touch each other, but of those which are separated by the exciting liquid, whether held in cells or interposed cloth. Erroneously ascribing the power of his series to an electromotive power, and conceiving the agent intervening between the metallic couple as acting merely as a conductor, Volta constructed his pile, as represented by fig. 10, in which there are two useless plates, one of each metal, at the extremities. The series really commences with the lowermost zinc plate, and ends with the uppermost copper plate, and as the current is from zinc to copper within, and from copper to zinc without, as represented by the arrows, fig. 10, it follows, that when rightly constructed, the copper end is positive, the zinc end negative. The addition, by Volta and his imitators, of two useless plates, as above stated, has led to a general impression, that the true zinc end of a series is positive, and the true copper end, of course, negative, contrary to the real state of the case.

355. I consider the original pile of Volta, and every other modification of the voltaic battery, as consisting of a series of elementary batteries, each alone competent to supply a galvanic current, and to the accumulated powers of which the energy of the whole is due. But such batteries do not severally consist of the two plates of metal which are in contact, or united by a wire or strap, but of two plates, forming a triad with the interposed exciting liquid. But if such a pile consists of such triads, making the base negative, we must begin the pile with zinc, and of course terminate it with

copper.

5

356. Since, when a wire or strap is made to proceed into a liquid, from the copper plate really forming the positive or delivering end of the series, it occupies the place which a zinc plate would occupy in the next additional cell in the couronne des tasses, or Babbington's apparatus, were the series extended, Daniell designates the surface of such a wire or strap, as the zincode." This appears to me objectionable, since it tends to produce a narrow association of a general property with a particular metal. Though it may be truly alleged, that the zincode occupies the place which the generating metal zinc, would occupy; yet, nevertheless, it does not act as a generating metal. It does not, like the preceding zinc plate, contribute to the current.

357. Faraday has proposed to call the poles of the voltaic series, electrodes, from 666, a way, and electron; the sectional polar areas thus designated, being the passages through which the electricity is assumed to flow into or out of the series when the circuit is in operation. Supposing the wires of the series, those of fig 9, for instance, to be introduced into water,

atoms of water which touches the wire proceeding from the copper; and the liquid, is called by the same author the anode, while the other mid the row in contact with the wire, proceeding from the first zinc pure at

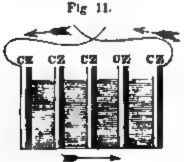
series, is called the cathode.

258. In order to understand the etymology of these words, it may be expedient to refer to the fact already mentioned, that the magnetic refer to the fact already mentioned, that the magnetic refer to the fact already mentioned, that the magnetic refer to the allege always to take a position at right angles to an electrical correct the influence of which it is sufficiently exposed. The direction of the pass needle is ascribed to such a current, caused by the progresses of solar heat from east to west, consequent to the diurnal rotation of the influence of any portion of the crust of the earth subjected to the death, or where the sun rises, hence anode, from sow, upwards, and way. Again, the fluid passing out westward, we have cathods for way. Again, the fluid passing out westward, we have cathods for solar with the ends of the wires forming the electrodes of a galvance beautiful to the part in contact with the positive pole will be the anode, while the pain contact with the negative pole will be the cathode.

359. I shall, however, continue to designate the poles as they have been heretofore named, using also Faraday's new appellation (electron) is either. Noad has judiciously, as I think, proposed that the positive pole designated as the anelectrode, the negative pole as the cathelectrode.

by Faraday, designated electrolysis, from electron, and hu, to untail free from bondage. A substance, susceptible of this process, is designated an electrolyte, and is said, when subjected to it, to be electrolysed. It mentary constituents are called tons, from io, to go. That which is evolved at the anode, anelectrode or positive pole, is called an anion, while the other, which is evolved from the cathode, cathelectrode or negative put.

the size of the pairs, and the whole number associated in the size of the pairs, and the whole number associated in the size of the pairs, and the whole number associated in the size of there must be two plates for every triad, this mode of estimation case to error in fact, however it may tend, when unnoaced, to be produce to hypothetical misconception. In the voltage file, fig. 10, the control to and electro-positive metals, copper and zine for instance, are in the copper of one triad to discharge to the zine of the other, as in the series, or triads, forming the series, consist each of a copper and zine place which do not touch or communicate otherwise than through the liquid to of those which are in contact as in the pile, or connected by a wire as the couronne des tasses, figure 9.



362. Soon after the pile and the courone of tasses were invented by Volta, another form of the voltaic series was contrived by Cruikshank, or of the most distinguished chemists of his time. Of this, figure 11 is a representation; usually it is a signated as the Cruikshank trough. The pair of this apparatus consist of a zinc and copper plat of similar dimensions, united so as to bring their corresponding surfaces in contact throughout. Each

pair forms one partition, consisting, on one side, of copper, on the other, of zinc. The partitions thus formed being inserted in an horizontal trough, have between them interstitial cavities for holding the acids, and which consequently perform the office of the cloth in the voltaic pile. Of course, the metals must be in the same alternate order as in the pile. The right side of the partitions being all of one kind of metal, those on the left must all be of the other: so that looking down at the trough obliquely from one end, we see none but copper surfaces; when looking from the other end, we see none but zinc. It is better, however, to have an extra copper plate at each end, to which to solder the rods or wires employed to communicate the galvanic current, or to act as the electrodes. These plates should be attached severally to the pieces of board forming the ends of the trough. This successive alternation of copper and zinc, or any other masses having a similar efficacy, whether organic or inorganic, I shall designate as voltaic order, being that which is essential to any modification of what we call the voltaic series, whether natural or artificial.

363. The analogy between the Cruikshank trough and the pile must be evident, since, if we place the trough upright, and after filling the cells between the pairs with moistened cloth, remove the wood, we shall have the

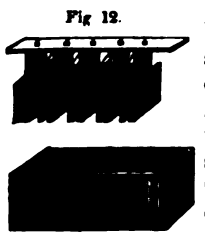
voltaic pile in the form of a rectangular prism.

364. The trough of Cruikshank has great advantages over the pile, or the couronne des tasses. As the energy of the current diminishes with the activity of the acid, and this is lessened in abundance and strength by reaction with the zinc, the quantity which can be held by the cloth is soon exhausted; and although, in the couronne des tasses, the supply of acid was as much too great, as in the pile it was too small, the plates could not be kept parallel, nor sufficiently near without liability to touch each other. In the trough there was an abundance of acid, and the plates, however approximated, were secure from contact: moreover, the Cruikshank trough, as since modified by me, in order to comprise an equivalent degree of efficacy, need not occupy a tenth part of the space required by the couronne des tasses. Again, the contact is immediate and more effectual between the zinc of one pair, and the copper of another.

365. The direction of the current in the trough is indicated by the arrows in the figure. It will be observed, to obey the general law of the circulation, as already mentioned, it proceeds from zinc to copper within the

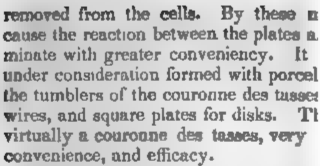
liquid, and from copper to zinc through the wire.

366. The analogy between Babington's apparatus, fig. 12, and the couronne des tasses, is as great as that which has been shown to exist between the trough of Cruikshank and the pile of Volta; the superiority of the apparatus of Babington over the couronne des tasses, being equally striking.



367. In Babington's apparatus, a trough of porcelain is so constructed, as to have in it partitions of the same material, dividing the interior into ten or more cells of a rectangular shape. With a view to economy both of the sheet metal employed and space, the plates are made also of a corresponding rectangular shape. We have seen that in the pile, and in the Cruikshank trough, the triads communicate by direct contact. In the apparatus of Babington, a communication between them is established by straps of cop-

per, which serve also to attach severally to one beam each couple appertaining to one trough, so that they may be simultaneously introduced into or



368. The great apparatus of Davy troughs of ten pairs each, upon Babing

two thousand triads.

369. Subsequently, agreeably to the paratus, otherwise constructed like th rounded by copper, contact being preve of the zinc was conceived to be increased.

370. The celebrated apparatus of Claston's plan; it consisted of twenty p copper coupled by a metallic strap, ar each pair was situated between the two sured six feet by two feet eight inches.

371. Subsequently, adopting the san respects surrounding the zinc by copper dred pairs, in which the vertical edge copper, instead of the upper and lowe importance in diminishing the necessithus constructed were separated by venot insulated any further than as copplarity on which voltaic action depends the arrangement of my apparatus, was throw the whole of the acid on or off of

The opinions of Faraday respecting the be found in the following abstract republic Sciences. They are preceded by the following suished editor of that Journal. See Vol. X.

"An account was given in this Journal (lume 3) by Dr. Hare, of his galvanic deflagration to their size in producing intense ignit Dr. Hare were, with some additions, repeate the adegations of the inventor were also Hare's memoirs and ours, with engravings, sophy and Philosophical Magazine, London, tors were imitated in Europe the old and in ral use. It is satisfactory to find, from the Faraday, that the course of his investigation deflagrator of our countryman as the most equote those passages of Mr. Faraday's paper tend to justify our statement.

"The advantages of this form of trough a exceedingly compact, for one hundred pairs more than three lect in length—2d. By Dr. upon copper pivots which rest up ne opper b tions, and these I have found it very converency, fastened in the front of the stind of the give the great advantage of arranging an appearance before the latter is put into action—3 use in an instant, a single jug of dilute acid hundred pairs of four inch plates—4th. On i

372. It was the power of simultaneously lowering or lifting all the plates attached to a beam, that caused a preference to be given to the apparatus of Babington's construction, over that of Cruikshank. Yet as it was found expedient, by Davy and others, to attach no more than ten couples to a beam, the object of an instantaneous and simultaneous immersion of the whole series in the liquid, or that of its extrication therefrom, could not be

effected conveniently upon the Babington plan.

373. I succeeded in this object by various constructions, which I have designated generally as galvanic deflagrators. It may be inquired, in what respect do series thus constructed so much differ from others as to deserve a peculiar name? The reply is, that in voltaic series, as employed by its inventor, and as used by all subsequent operators, the plates were necessarily subjected to the exciting reagent in the first instance, the circuit being completed by a communication, subsequently made, between the electrodes. In that construction, to which I give the generic term deflagrator, the circuit may be completed by the simultaneous exposure of the whole series of plates to the exciting reagent, either simultaneously with, or subsequently to, the completion of the rest of the circuit.

374. Turner alleges, that an advantageous variation of this contrivance was made by Mr. Hart, of Glasgow, by which the copper was formed into a hollow parallelopiped. This construction was resorted to by me as early as the year 1822, but not finding it as convenient for filling and emptying

ter of a revolution, it becomes active, and the great advantage is obtained of procuring for the experiment the effect of the first contact of the zinc and acid, which is twice or sometimes even thrice that which the battery can produce a minute or two after. 5th. When the experiment is completed, the acid can be at once poured from between the plates, so that the battery is never left to waste during an unconnected state of its extremities; the acid is not unnecessarily exhausted; the zinc is not uselessly consumed; and, besides avoiding these evils, the charge is mixed and rendered uniform, which produces a great and good result; and, upon proceeding to a second experiment, the important effect of first contact is again obtained. 6th. The saving of zinc is very great. It is not merely that, whilst in action, the zinc performs more voltaic duty, but all the destruction which takes place with the ordinary forms of battery between the experiments is prevented. This saving is of such extent, that I estimate the zinc in the new form of battery to be thrice as effective as that in the ordinary form. 7th The importance of this saving of metal is not merely that the value of the zinc is saved, but that the battery is much lighter and more manageable; and also that the surfaces of the zinc and copper plates may be brought much nearer to each other when the battery is constructed, and remain so until it is worn out: the latter is a very important advantage. 8th. Again, as, in consequence of the saving, thinner plates will perform the duty of thick ones, rolled zinc may be used; and I have found rolled zinc superior to cast zinc in action; a superiority which I incline to attribute to its greater purity. 9th. Another advantage is obtained in the economy of the acid used, which is proportionate to the diminution of the zinc dissolved. 10th. The acid also is more easily exhausted, and is in such small quantity that there is never any occasion to return an old charge into use. Such old acid, whilst out of use, often dissolves portions of copper from the black flocculi usually mingled with it, which are derived from the zinc: now any portion of copper in solution in the charge does great harm, because, by the local action of the acid and zinc, it tends to precipitate upon the latter, and diminish its voltaic officacy. 11th. By using a due mixture of nitric and sulphuric acid for the charge, no gas is evolved from the troughs; so that a battery of several hundred pairs of plates may, without inconvenience, be close to the experimenter. 12th. If, during a series of experiments, the acid becomes exhausted, it can be withdrawn, and replaced by other acid with the utmost facility; and after the experiments are concluded, the great advantage of easily washing the plates is at command. And it appears to me, that in place of making, under different circumstances, mutual sacrifices of comfeet. power, and economy, to obtain a desired end, all are at once obtained by Dr. Hare's form of trough."

as other constructions, the parallelopipeds were laid aside, and are calls

my laboratory.

375. I shall introduce into an appendix, engravings and descriptors several forms of voltaic series to which I successively resorted. The settlement to which I have finally given preference, is that of a Creation trough, so modified as to increase its power, to enable it to act as a designator, and render it possible to scrape the plates.

\* Englaving and Description of a Deflagrator of Two Hundred Pairs of Inches by Seven and a half, being one of the Members of a Pair of Troughs, in 4 the same size, and which may be used collaterally or consecutively.



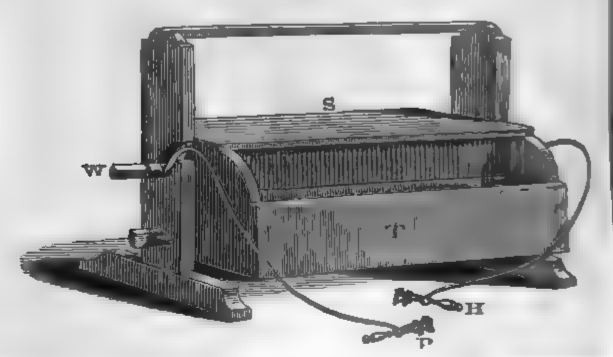


Fig 13 represents a deflagrator. It consists of a wooden trough upon the Cont. shank plan, joined lengthwise, edge to edge, to another trough so that be sides of the one are vertical, those of the observations be harizontal The a THAT of this arrangement is, that by a partial revolution of both the troughs, thus ar is upon pivots which support them at the ends, any fluid which may be in one trough must flow into the other; and, reversing the movement, must flow back again. T> galvanic series being placed in one of the troughs, the acid in the other, by a more ment such as described, the plates may all be instantaneously subjected to the acor relieved from it. The pivots are made of iron, coated with brass or copper & less hable to exidezement. A metallic communication is made between the catter of the pivots, and the galvanic series within In order to produce a connexion be tween one trough of this description and another, it is only necessary to all wa next of one trough to revolve on one end, and a pivot of the other trough to revolve and the other end of a strap of sheet copper. To connect with the termination of an series, the leaden rods, to which are soldered the vices, or spring forceps, for heat & the substances to be exposed to the delligrating power, one end of each of the cat rods is soldered to a piece of sheet copper. The pieces of copper thus soldered by the lead rods, are then to be placed under the pivots, which are, of course, to be connected with the terminations of the series. The last mentioned connexion is correctly made by means of straps of copper, severally soldered to the pixots, and is poles of the series, and screwed together by a land-vice. Each pair consists of a copper and a zinc plate, soldered together at the upper edge, where the coppera-made to embrace the edge of the zinc. The three remaining edges are made to exist For each inch in the length of the trough there are three a groove in the wood pairs. In the series represented by fig. 13, there are one hundred pairs of near t fourteen by seven and a half inches

The pair of troughs, forming, when associated, a single deflagrator, of which essis represented by fig. 13, may be employed either by connecting them at their home.

# If the different Modes in which the Power of a Galvanic Battery may be augmented.

376. Obviously there were two modes in which the ower of a galvanic battery could be augmented; one was hat of increasing the size of the plates of each pair, the ther that of increasing the number associated as a seies.

377. It has been mentioned, that under the name of caprimotor, the elementary battery or galvanic triad (312) as first made of gigantic size in this country (324). By Vollaston it had been tried in the minute form of a silver himble, of which the cavity was occupied by a minute ylinder of zinc surrounded by a potent exciting liquid. In the calorimotor the number was reduced to unity, while the size was enormously increased, the liquid being ighly energetic. In Davy's pile of two thousand pairs, he number was comparatively very large, the energy of he liquid as great as could be used with advantage. By assiot, the number has been carried to three thousand we hundred and twenty, excited by a liquid of no higher nergy than river water.

378. In the voltaic series thus employed by Crosse and Lassiot, we see that the number of the triads is exceedagly large, while the activity of the liquid is brought down

eneous or heterogeneous extremities. When connected at their homogeneous ends, ney act collaterally as a series of one hundred pairs of fifteen by fourteen; when onnected at their heterogeneous extremities, they act consecutively as a series of we hundred pairs, each fourteen by seven and a half. Of course, agreeably to the ret mentioned arrangement, we double the quantity, the intensity being unaltered: coording to the arrangement last mentioned, the intensity is doubled without an acrease of the quantity.

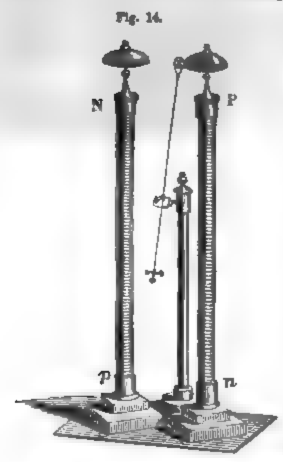
It may be proper to mention, that these Cruikshank deflagrators are so contructed, as that the plates may be scraped after each operation. Every fifth plate is emented, so as to form a water-tight partition. As the plates are purposely so cut, s to lessen in width from the upper to the lower edge, those which are not comented may be easily withdrawn in order to be scraped. But the withdrawal of every four moveable plates, leaves vacant spaces sufficiently large for the introduction of a tool

nitable for scraping those which are not removable.

A battery of this construction was made, under my direction, for the Lowell Instinte, consisting of four troughs, each containing one hundred plates six and a half
nches square. A piece of wooden plank, three inches wide, thirteen inches long,
nd two inches in thickness, is so bored by a centre-bit as to have eight cylindrical
avities, all these being supplied with mercury. By means of ropes of copper wire,
he various portions of mercury are made to communicate each with one of the eight
oles of the four troughs. Arches of stout copper wire are provided of different
angths, so that, by variously connecting the mercury in the holes, the whole may
e made to cooperate as one series of four hundred pairs each of forty-two inches,
ne series of two hundred pairs of eighty-four inches, or one series of one hundred
airs of one hundred and sixty-eight inches surface.

to a minimum, even below that of the saliva in the evenment with the tongue (306), or the sanguineous slime of the frog in the experiment of Galvani (309). But in the column of De Luc no liquid reagent is employed, suce t cannot be pretended that the hygrometric moisture of the paper, entering into its construction, exists in the land form. This series, as originally contrived, consisted of disks of silvered paper alternated with disks of zinc as the as possible, and each of about five-eighths of an inch a diameter. An analogous series, devised by Zamboni, consists of disks of the same size, coated on one side with left tin, on the other with a paste of black oxide of manganess and sulphate of zinc. According to Daniell, a senes of this last mentioned construction has been made, comprising 20,000 pairs, respecting the powers of which account will be given in due course.

Engraving and Description of the Electric Columnar Apparatus of R Luc, as modified by Zamboni.



mentioned construction, the adjoint figure is a representation. It counts of two columns, which communicate by a plate of zinc. The letters P p show the ends forming the poster poles, while N n indicate those wroing the negative poles. The harmone, and the cathode of the harmone municating through the remaining through the remai

380. No doubt the electric starts owe their efficiency to that the rate metive process, which, in other a, or causes an electric accounting to be electrodes; but the phonomenant most exactly those of electricity cannot by friction.

381. When the apparatus of a sented by the engraving was root a constructed, it produced sparks at the

contact of the bells with the pendulum. It communicated a charge is small coated glass cylinder, sufficient, in passing through the thumbard finger, to give a shock extending to the wrist. After it had been emstructed for more than ten years, when warmed it would cause a gold on to vibrate between balls severally communicating with the electrodes. The

the columns comprise about 4500 disks of tinned paper, coated, on the stherwise naked side, with manganese and sulphate of zinc.\*

On the Construction of De Luc's Columns, as modified by Zamboni; and on a Modification of the Single Lauf Electrometer contrived by the Author, by which the ultimate Eclency of a large electric series may be ascertained, by testing a small portion of the members of which it is to be constituted.

. About fifteen years ago, the construction of De Luc's electric columns, as modi-fied by Zemboni, was undertaken by Isaiah Lukens, one of our most skilful and innieus mechanicians.

The materials employed were paper covered with leaf tin (erroneously called sil-

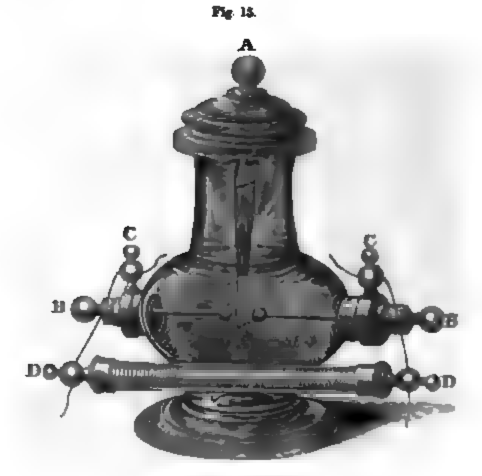
The materials employed were paper covered with lear un (erroneously cause materials appear), peroxide of manganese, and crystallized sulphate of sinc.

The peroxide was finely pulverized and mixed with a concentrated solution of the sulphate. The mixture thus formed was, by means of a brush, applied like a pigment to the surfaces of the paper not coated by the tin. The sheets were afterwards appear out on the floor of an apartment, and left during the night to dry. By these means, unnecessary exposure to light was avoided, which Mr. Lukens conceives to the injurious, especially as received directly from the sun. Next day the sheets were est into disks of about five-eighths inch diameter, by means of a hollow punch. The blake were then niled, with the heterogeneous surfaces alternating, as in other voltain. disks were then piled, with the heterogeneous surfaces alternating, as in other voltais paries, and were introduced into, and compressed within, glass tubes, accounted as usual with pedestals, caps, and bells. Notwithstanding his skill and experience, Mr. Lukens latterly complained of occasional want of success, arising, as he supposed, from the defective quality of the manganese. In various instances, his columns, after being constructed with the utmost care, proved inert.

The manipulation, likewise, according to his plan of operating, appeared to me to be troublesome and precarious. He was accustomed to place a row of the disks, as large as could be conveniently handled, in a trough of sheet metal, and then transfer the pile thus formed to the glass tubes. This operation, to be successful, required

dexterity.

In February, 1833, wishing to replenish the tubes of a pair of electric columns which had become effets, I contrived to avoid the risk of expending the labour and attention requisite to finish a series, while uncertain as to its eventual efficiency. I



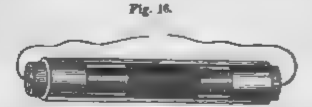
#### Of Groves' Gas Battery.

382. "In the L. and E. Phil. Mag. for December, 1842, a most runs dinary and perfectly novel voltaic battery is described by the invente. It

contrived, likewise, by a very simple expedient, to facilitate the process of play to

The first mentioned desideratum was obtained by means of a single less of suspended in a glass vessel (represented by the subjoined, figure), between the knobs at the ends severally of two brass rode, B B, proceeding through repeated of the vessel towards each other, so as to be capable, if requisite, of meeting by cantre. By means of screws, the knobs on these rods were susceptible is adjusted to any distance from the gold leaf suspended between them. Lateral leaf thus situated, vibrations may be produced by a series of disks, comprast more than one-twentieth of the number necessary to cause such a pendular at monly pertains to the electric column, to oscillate. In the case in point is that the disks produced by one sheet of paper, were sufficient to make the set of brate setively between the knobs. The mode in which this effect was pressed, in due order, within a glass tube, by spirals of wire.

Each of the wires of which these spirals were formed at the ends enclosed at tube, being unaltered throughout the remaining portions of their length, was post-through corks closing the crifices of the tube. The series thus prepared, as the series of the tube.



placed in the situation of the electric column, appended to the instrument agreed to fig. 15, being in like manner suspended from the rods outside of the result means of the projecting wires already mentioned. Thus situated, if there by adequate degree of electromotive power in the series under trial, and the structure sufficiently dry, the excitement of the poles will be communicated to the holes be indicated by the consequent vibrations of the gold leaf suspended with

When a larger series is used, such as that represented at D D first with the discontinued, only in consequence of the adherence of the adherence of the adherence of the highest the highest built at the vertex of the instrument, to which the force of the leaf is affixed. The finger being removed vibratory pulsations will recommend be sooner or later arrested in the same imaner as at first

When the knobs BB, were properly connected with the poles of a vitar town of seven hundred pairs excited merely by pure water the pulsations of the cut of quick and incessant. These pulsations may incente the chettre interest and not furnish any criterion of the divellent igniting, or electro-magnetic pieces to voltate series.

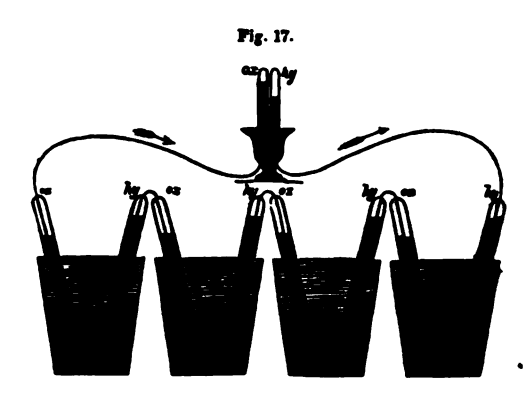
It may readily be perceived, that the electrometer, constructed as been a ceribed, constitutes an electrical discriminator, which may enable up to the electrometive powers of various substances arranged as disks in a series of a lings to disks. I have already ascertained that aurum musicum, spread of the timed paper, produces an electrometive series.

The piling of the disks was facilitated by using a punch excavated so at a man a point in the centre, by which the centre of each disk was put tured. It are of the puncture thus made, it was easy, even for an unaxidial persists and them concentrically upon a silk thread, and to transfer them to the turns of derangement.

The manganese which I employed with success, in the replemation of the or tric columns alluded to above, consisted mainly of needle shape i rate of the aggregated into lumps. Mr Lakens alleges that the crystal and manganese always, agreeously to his experience, proved the best for the construction of employment.

The electrometer, with an electric column attached to it, as above representation

Frove. It consists of a series of fifty pairs of platinized platinum plates, ach about a quarter of an inch wide, enclosed in tubes partially filled, al-



ternately with oxygen and hydrogen gases, as shown in fig. 17. The tubes being charged with dilute sulphuric acid, sp. gr. 1.2, the following effects were produced:—

"1st. A shock was given which could be felt by five persons joining.

hands, and which, when taken by a single person, was painful.

"2d. The needle of a galvanometer was whirled round, and stood at about 60°; with one person interposed in the circuit it stood at 40°, and was slightly deflected when two were interposed.

44 3d. A brilliant spark, visible in broad daylight, was given between

charcoal points.

3

3:

is E.

"4th. Iodide of potassium, hydrochloric acid, and water acidulated with sulphuric acid, were severally decomposed: the gaseous elements from the decomposed water were extricated in sufficient quantity to be collected and detonated. They were evolved, as chemical theory and experience would indicate, the hydrogen at the anode, the oxygen at the cathode. It was

may serve to show the nature, as well as the extent of electric excitement; since, when an electrified mass is made to communicate with the brass ball, A, from which the leaf is suspended, the latter ought to be attracted by that knob, which receives from the series an opposite excitement. Hence, the excitement of the electrified body being known, that of the poles may be detected: or the latter being known, the excitement of the body may be discovered. This application of the electric series is not, however, a new idea. I saw many years ago a notice of an electrometer, associated with an electric column, in such manner as to be used as an electrical discriminator.

The great difficulty in resorting to this means of discrimination is, that an electrified body may, by induction, produce in a conductor alternately, opposite states of electrical excitement. As it approaches the conductor, it may cause it to receive, or give out electricity; of course, when retracted, the conductor will have the opposite excitement to that consequent to approximation. Supposing the brase ball of the electrometer in question, to be affected in the mode just described, the leaf suspended from it must be successively attracted by each pole. Besides, the excitement may be so strong, as to render that of the scries nugatory; as in the case of a powerful magnet, which will attract either pole of a feeble one.

The direction of the first pulsation of the leaf is the best criterion; but reliance should not be accorded to one experiment, especially when so easily repeated. I find that a gilt pith ball, if suspended in place of the leaf, will vibrate for a time. It is, however, like the leaf, liable to have its movements arrested by an adherence to

one or other of the knobs.

found, that to decompose water, twenty-six pairs were requ four pairs would decompose iodide of potassium.

"5th. A gold leaf electroscope was perceptibly affected.

888. "When the tubes were charged with atmospheric air, so dista produced, nor was any current determined when the gases employed w

carbonic acid and nitrogen, or oxygen and nitrogen.

384. "The original experiment was not made with sufficient ends to show with accuracy the proportional diminution of gas in each se though it was plain that the hydrogen diminished much more rapids the oxygen."

385. The electrodes are represented as terminating in tubes for cati the gaseous elements of water, evolved by the electrolytic power of! same elements, under the electro-polarizing influence of platinum.

#### Of the Electrolytic Process by which the Galvano-Electric Curve propagated within Liquid Reagents.

886. Before giving an account of these forms of the galvanic series which more than one chemical reagent is employed, it may be expedien enter more fully into the rationale of that part of the galvanic promi which the electric fluid is supposed to be transferred from the zinc to

copper, through the exciting liquid.

387. Faraday assumes, that the fundamental cause of the hydrocia cal current, or that produced by the reaction of metals with liquids, at mical affinity between one of the metals employed and an ingreduct some electrolyte contained in the liquid, and that the combination atom of the metal with an ingredient atom, taken from the liquid elected causes an equivalent discharge of electricity from the atom so taken.

388. Thus, when zinc combines with the oxygen of water, perfort the part of an electrolyte, a discharge of electricity from the atom of t gen is a necessary consequence. Hence, in some cases, as, for the that of pure zinc exposed to diluted sulphuric acid, unless the process charge be facilitated by contact with a metal not capable of combining oxygen, no combination ensues. But, under the same circumstances while the oxygen of the liquid is attracted by zinc, the liquid be in or with a plate not liable to be oxidized by it, the discharge of electricity a cilitated, as soon as a conducting communication is established between metals. He supposes a series of decompositions and recompositions to place within a row of the particles of water lying between the zinc and per surfaces, so that the atom of oxygen next the zinc surface, unling an atom of this metal, discharges its electricity to the next atom of by gen, which unites with the oxygen of the next atom of water. But n 2 this, it causes the transfer of its electricity to the third atom of helic This process being repeated, an atom of hydrogen surcharged with ee city, is evolved against the copper surface. Hence the surcharge acc. by the copper goes through the connecting wire, to restore the ext tension within the zine which the chemical reaction tends to dim as above stated.\*

<sup>&</sup>quot; If we suppose, in a dance, a number of girls and hoys standing in a row, in ples, alternately arm in arm, alternately separate, and that a girl is suddenly from one end of the row, and a boy from the other end, evidently of each extwould be an individual isolated. But if, simultaneously, each girl relinquishia hold on the partner of which she had the arm previously, takes that of the nex on the other side, there will be no individual of either sex in a solitary state

er 389. The id a t se of hy rogen at one electrode, and the i of decompositions and recommygen at the other, is the effect of a exositions, as above stated, was or ted by Grothus, as account-1 ang for the fact, that the extrication or hydr n and oxygen from electroysed water does not take place at t It had been objected )t. 10 that it was inconceivable that the , arising from the deents of w composition of a succession of atoms of this liquid, could be severally avolved at the distance of some inches from each other.

390. In the last edition of my Treatise on Galvanism, after having cited the electrolytic theory of Faraday, (386, 387,) I subjoined the following

remarks:-

391. Whatever may be the source of the imponderable matter which circulates in voltaic series, I infer, that its confinement within the circuit, and its endowment with an ability to pass over an interval, or to penetrate, through imperfect conductors, must be due to an influence of the masses

concerned, which has not as yet been satisfactorily explained.

392. Probably upon an analogous ability to produce or annul, to promote or retard, chemical reaction, the efficacy of animal and vegetable organization is founded, being obviously dependent on an arrangement of masses. The voltaic series of a gymnotus is evidently an animal organ, and its analogy with the voltaic series, produced by human ingenuity, induces me to consider the latter in the same class of agents as the organs by

which life is supported.

393. The opinions thus expressed by me have, I think, found much to justify them in the facts established by means of Groves' gas battery. In that apparatus, a portion of hydrogen performs the part of zinc, while oxygen performs the part of copper (314). These gases exercise chemical affinity neither with water nor with platinum. An electro-polarizing reaction with the platinum is evidently the only imaginable cause for the union of an atom of gaseous hydrogen with the oxygen of water, at one of the terminating surfaces of that liquid, while, analogously, an atom of gaseous oxygen combines with hydrogen at the other aqueous surface.

394. It appears to me, that precisely the same change must ensue in the electro-polarity of the gaseous atoms during this electrolytic process, as when a mixture of the gases employed, is made to combine by the presence

of platinum sponge.

395. But whatever may be the rationale, the phenomena of this gas battery, as well as those previously used, all tend to confirm the fact, that what is called the electric current, in its passage in a galvanic triad from the electropositive surface to the electronegative surface through the liquid, is inseparably associated with a series of decompositions and recompositions.

396. This electrolytic exchange of partners, in rows of atoms which, within the battery, appear to be concerned in the generation of the current, seems, externally, to be one of its effects; so that the evolution of the elements of an electrolyte, severally at the cathode and the anode (358), are ascribed to a series of decompositions and recompositions, like those which

accompany the generation of the current.

397. The impression seems to exist among many electricians, judging from their language, that the elements actually travel from one electrode to the other. The choice of the word ion, from io, to go, (360) is in this respect disadvantageous. In point of fact there can be no such locomotion. Simultaneously with the liberation of an atom of each ingredient at the poles, the residual ions in the electrolytic row, rearrange themselves so as

of the electrolytic atoms will be simultaneously decomposed, out of a major rows existing in due contact with the electrodes; but it cannot be said ably supposed, that there is not a change of relative position too scenario permit the electrolytic process to take place twice through process to take place twice through process to the interposition of precisely the loss of an atom can never and the interposition of precisely the same residual atoms which have can once together in a row long enough to lose one atom from among the As reasonably might a sportsman expect to find in a numerous had a birds while wheeling about him, the same individuals twice in a line with

rested fowling piece.

398. Between the galvanic batteries above described, in which out a liquid is employed, and those constructed with the aid of two hours des cal reagents or electrolytes, as in the case of the series of Daniel or Game. there is this striking difference, that, in the requisite series of decession tions and recompositions, the part performed by hydrogen as the compositive element on one side of an interposed disphragm, is on the dist # formed by copper, nitric oxide, or some other radical more ready that it drogen to separate from oxygen. The porous diaphragm soaked water phate of copper or nitric acid, presents throughout the whole of its are oxidizing surface which cooperates with the opposite influence of the oxidizing zinc to effect the disunion of the elements of water. Hence the electrolytic exchange is solicited on one side, by the attraction of rint in exygen, it is solicited on the other side by the attraction between by ing and the oxygen of the other electrolyte with which the pores of the duplete. are imbued. In nitric acid, nitric oxide, acting as a compound rade at ... forms the part of the simple radical copper, in the oxide of this metal 🚟 in the sulphate. The liberation of one atom of nitric oxide, at the surface of the platina, causes three atoms of oxygen to be yielded to the hydrogen of water at the membrane, more readily than an atom of copper, and gously liberated, causes one atom of oxygen to be yielded. That come ceive to be the main cause of the superior efficiety of Gross's a terms. sustaining battery, of which a description is given in the following pur-To the advantage thus gained is to be added that of the protection of # metallic surfaces from injurious deposition (315).

#### Of Constant or Sustaining Batteries.

London, made an interesting and important innovation in the construction of galvanic batteries, by the interposition of an animal membrane, the gullet of an ox for instance between the heterogeneous metallic surfaces. A battery of this kind in my possession, consists of a cylinder of copper, about six inches in height and four inches is diameter. Within an inch and a half of the brim, a diaphragm of copper is inserted. This diaphragm is perforated like a colander, and has, at the centre, a hole of about an inch and a half in diameter. It is through this hole that the gullet is made to descend to the bottom of

the vessel, where it communicates with a glass tube, which is so bent at right angles twice, and so extended as to rise by the side of the cylinder and terminate in a recurved orifice on a level with the brim. A solid cylinder of amalgamated zinc, of half an inch in diameter, is made to oc-cupy the cavity of the gullet.

400. It has already been mentioned, that in this construction, the copper is not liable to be tarnished nor the zinc to be coated by a deposition,\* and thus to be rendered incapable of their appropriate reaction with the interposed liquid. The membrane will not allow a solution of either metal to permeate it, although it permits the transmission of the electro-chemical decomposition and recomposition, by which the voltaic current is sustained. The space between the gullet and the cupreous surface of the enclosing cylinder, is occupied by a saturated solution of sulphate of copper, the cavity within the gullet by diluted sulphuric acid, or some saline solution not liable to be decomposed, per se, by zinc. Upon the diaphragm, crystals of sulphate of copper are deposited, for the purpose of preventing any decline in the strength of the solution of that salt. As the diluted acid employed becomes saturated with zinc, it is displaced by a fresh portion which causes that first mentioned to be expelled at the orifice of the glass tube. Of course, the chemical reaction may be kept up as long as the zinc lasts. Latterly, in the construction of Daniell's battery, the glass tube has been omitted, as the advantage arising from its employment, has not been found to compensate for the additional liability to derangement.

401. I saw at the laboratory of Prof. Daniell, an association of his sustaining batteries, of which the power to

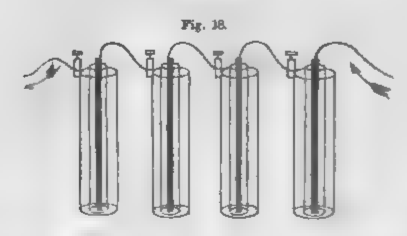
ignite wire was surprising.

402. According to Brande, with a constant battery of twenty cells arranged in a single series, twelve cubic inches of mixed hydrogen and oxygen may be collected in every five minutes of action; and, when they are connected in pairs and afterwards in a series of ten, the quantity amounts to seventeen cubic inches. Eight inches of platinum wire, one two-hundredths of an inch in diameter, may be kept permanently hot by the same arrangement, and the spark

The deposition thus formed consists of a mixture of copper and suboxide of zinc (315, and note).

taken between charcoal points is very brilliant. To me original form of Daniell's battery above described, there however, a great objection, founded on the employment of an animal membrane, which is of course liable to be a jured or spoiled. Porous vessels of unglazed eartheaver have been, however, substituted for animal membrane with great increase of convenience, and they are an employed both in this and other forms of voltaic appearance. Leather has also been successfully substituted for membrane.

Engraving and Description of a Voltaic Series of Constant Batteries.



403. The preceding figure will convey a sufficient conception of a sustaining battery, in which cylinders of porous porcelain are substituted for the ox gullet employed agreeably to the original construction of Daniell's ap-

paratus.

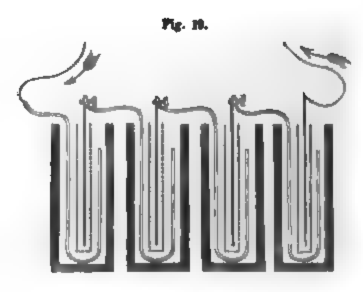
404. The external lines represent sections of round coper jars, the dark central lines represent the amalgamated zinc rods, situated in the common axis of the copper jars and of the cylinders of porous porcelain. These are represented as occupying a mean position between the unknown and copper. The charge of the apparatus and rationar of its operation, is the same as that already stated in reference to the constant battery in its original form.

405. I do not believe the effects of any apparatus constructed upon this plan, to be at the outset as powerful striking as those which are producible by a deflagratore a like extent of surface. Still less would it be possible to comprise a series of like degree of power within a space equally small. A series of two hundred pairs of sustaining batteries would be extremely unwieldy and cumbrous. It

such, as to enable me to withdraw four plates out of any five, so as to have the whole accessible to a scraper. Preover, the amalgamation of the zinc surfaces, devised Sturgeon, confers upon the plates of the Cruikshank astruction, to a certain extent, the properties of a susning battery. The great objection, to the somewhat ally use of mercury, is, that it enfeebles the zinc plates so to make them liable to fracture; but this is less to be prehended when the zinc is supported by corresponding oper plates, as in pairs of the kind above designated, and ere the mercury, as in these pairs, is to be applied only one of the surfaces of each zinc plate.

## Of Grove's Sustaining Battery.

106. About the year 1840, Prof. Grove made a very insting modification of Daniell's battery, not only by stituting vessels of porous porcelain for the ox gullet, also by using platinum and nitric acid in lieu of copper I a solution of the sulphate of that metal. The porous, nearly full of nitric acid, is surrounded by zinc instead in sulphuric acid diluted with only four parts of ter, contained in a sufficiently capacious receptacles intric acid, by oxidizing the hydrogen, accelerates the atrolysis, nitric oxide is evolved at the surface of the tinum, which is made to occupy an intermediate position, within the porcelain cell.



107. The dark central lines (figure 19) represent secus of the platinum plates; the dotted lines C, which inde these, represent the hollow cylinders of porous porcelain. The double lines, which comprise both of the already mentioned, represent sections of the zinc operates. The dark external lines, in like manner, represent the glass jars within which the rest of the apparatus situated. The arrows indicate the course of the communication of

408. A series of this kind of only four pairs, constructed in my laboratory, caused the long continued ignitice about two inches of platinum wire. According to Prof. Grove, a series of one hundred pairs, each of the same a mensions as those above described, caused a dirupture for charge, accompanied with intense heat and light, through an interval of three inches, which is as much as was fected in this way, by the great battery of two thousand pairs with which Davy operated at the Royal Institutes.

409. Latterly it has been found that plumbage or col may be substituted for platinum in Grove's apparate This material was resorted to independently both by some German chemists, by Prof. B. Silliman, Jr. and myss An extensive series was constructed for the Lowell Inc. tute, by Prof. B. Silliman, which was very powerful at had the advantage over the deflagrator of a much me durable intensity of action. In practice, however, the are several objections to Grove's apparatus. The second introduction of sulphuric and nitric acid into a great number ber of receptacles in a much more concentrated state that that in which they are used for other voltaic series, as wias the subsequent removal, is a very troublesome and (= agreeable task. Moreover, the fumes of nitric oxide who are copiously evolved, are extremely annoying to all persons exposed to them, and are to some persons insured The porcelain cells are very liable to be broken and soon suffer by the disintegrating influence of the acids.

410. In an arrangement suggested by Berzelius, a cylinder of coke is hollowed out into the form of a cup, my which the nitric acid is placed, and thus the use of a earthenware diaphragm is avoided.

411. A great difficulty arises from the expensive neces

sity of employing platinum as the material of the screws and wires, by which the requisite attachments are made in batteries formed by the aid of carbon, as above described. Any other metal, excepting gold, which is three times as dear as platinum, is liable to be corroded by the nitric acid, which rises in the carbonaceous pores, by capillary attraction.

# Of the Passive Iron Battery.

412. The battery thus designated, was first suggested by Mr. Hawkins (L. and E. Phil. Mag. Vol. XVI.), in consequence of the previous discovery by Herschell, Schoenbien, and others, of the susceptibility of iron of what is called the passive state, which renders it insusceptible of being acted upon by strong nitric acid; whereas, in its ordinary state, it is oxidized and dissolved in that acid. The passive state does not protect the metal from acid of the s. g. 1.2. This state of iron is induced by the contact or association with platina, of which, being highly electronegative, the presence by analogy with copper and zinc should promote reaction with corroding reagents (321). The passive state also protects iron when exposed to diuted sulphuric or phosphoric acid, but cannot avail against shlorohydric acid.

413. This phenomenon is one of the most mysterious, put it is not expedient now to enter into any discussion respecting the cause. The object is to mention the use made of this wonderful property. As the inactive condition is spontaneously assumed when iron is plunged into acid of s. g. above 1.35, a plate of this metal has been substituted in a Grove's battery for the platina plate. According to the authors above named, batteries thus formed were found to be efficient; but, unfortunately, the peculiar condition of the iron is liable to cease. When this happen the rapid solution of the negative plate follows.

414. Schoenbien mentions the formation of batteries either of zinc and passive iron, or altogether of iron cylinders, one set being passive, the other in the normal condition. Five cast iron cylinders, 10 inches high, 3† diameter; as many of zinc of same height, and 3½ in diameter. Intermediate between the zinc and iron, porous cylinders of earthenware, one part nitric, s. g. 1.4, with from twelve to three parts sulphuric acid, were found to

answer. Subsequently, cylinders of iron, in its ordinal state, were substituted for zinc cylinders. By the tery, 2400 cubic inches of the gaseous elements of were liberated in one hour.

415. It seems, however, that the passive condition of iron is liable to cease, and when this happens, the metals dissolved. I am under the impression that butteres of this kind can never be found advantageous, as, independently of the uncertainty arising from the precariousness of the passive state of iron, this metal is very liable to be prize in jured by rust when out of use.

## Of Smee's Battery.

vered that the efficacy of a battery, made like calonal tors or deflagrators (324, 375), Babbington's apparatus (367), or one of his own contrivance, in which a thin she of silver is situated between two plates of zinc, may be augmented by a deposition of platina sponge being upon the surface of the negative metal. For this purpose platina, palladium, silver, or plated copper, may be used but, taking both efficacy and economy into view, silves.

rolled very thin, was found preferable.

417. In the case of a constant or sustaining batter. (398) it has been stated that the oxidizement of the hydrogen at the diaphragm, obviating the necessity for its relation in the gaseous form at the cathode, facilitates to electrolytic part of the electromotive process in a galvant circuit. In Smee's apparatus, the extrication of the hydrogen at the cathode, is alleged to be facilitated by the mechanical power of the asperities of the platinizing deposition. I infer, however, that the contact of the platinum may have a favourable polarizing influence on the hydrogenous cathion, promoting its separation from the other element of water, analogous to that exercised by the same metal in the gas battery (393-4).

418. The following directions are given for platning the plates: "Each piece of metal is to be placed in water to which a little dilute sulphuric acid, and nitro-muriate of platinum, is to be added. A simple current is then to be formed by zinc placed in a porous tube with dilute acid when, after the lapse of a short time, the metal will be

The trouble of this operation is most trifling, only requiring a little time after the arrangement of the apparatus, which takes even less than the description. The cost is about 6d. a plate, of 4 inches each way, or 32 inches of surface. It is necessary to make the surface of the silver rough, by brushing it over with a little strong nitric acid, which gives it instantly a frosted appearance, and after being washed, it is ready for the platinizing process; but the finely divided platinum does not adhere firmly to very smooth metals."

#### OF THE PHENOMENA OF GALVANISM.

Under this head we may put effects upon the animal organs, statical effects, ignition of continuous conductors, deflagration, electrolysis or decomposition, galvanic powers and effects of animal organs, magnetic influence or electromagnetism, galvanic effects of heat, or thermo-galvanism, sometimes called thermo-electricity, or thermo-magnetism.

#### EFFECTS UPON ANIMAL ORGANS.

Effects on the Organs of Taste, on those of Sight, Nerves of Sensation; in producing Convulsions.

419. Electricity is not the only one among the supposed imponderable elements, which can cause sensation. The influx and efflux of caloric produces a sensation in one case of cold, in the other of heat; moreover intense light, like that of the sun, of galvanic ignition, or the oxyhydrogen lime light, produces in the eyes a painful feeling. But in giving shocks to live animals, and in causing muscular movements or convulsions in a carcase or corpse, electricity has a peculiar attribute.

420. Probably the first knowledge of the shock produced from a galvanic circuit, arose from experiencing that given by the torpedo and other electrical fishes. This wonderful faculty could not but have been observed in the most ancient times by fishermen in the countries where such ani-

mals are found.

421. A very feeble discharge like that between a zinc and silver disk, including the tongue, has been felt from the

earliest period at which wine, cider, or beer of any kind was drunk from a metallic vessel. The preference great to pewter as the material of a mug from which to dreat beer, is now ascribed to a galvanic discharge arising from the metal, the lips and the beverage, forming a feeble givenic triad, in which the liquid is electropositive, being slightly acidulous. Probably from an analogous cause arises the characteristic taste of acids, while from an opposite galvanic discharge in which the solution is electropositive, arises the peculiar savour of the principal alkales. I am inclined to the idea that all the various tastes and odours, which we are liable to experience, are due to modifications of galvano-electro-chemical causes.

422. But the first recorded notice of the phenomena of a galvanic discharge, as respects its obvious production by a metallic triad, was published not as associated with a theory directed to the explanation of physical phenomena; but on the contrary, in a metaphysical effort to suggest a theory of pleasures. In the year 1767, more than twenty three years before the convulsions produced in frogs by metals was noticed by Galvani, Sultzer, the author of the theory of pleasures alluded to, gave an account of the taste produced by heterogeneous metals, when including the tongue, and brought into contact at their projecting

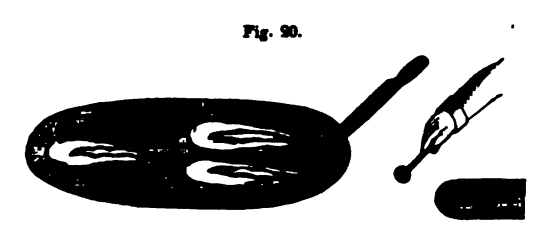
extremities.

423. It must be evident that the original observer of the phenomenon, thus described by Sultzer, was the discovered of the elementary galvanic battery; and it is remarkable that Galvani was led to infer the existence in nature of the powers of such a battery, by a phenomenon which had no connexion therewith. The hind parts of the carcases of some frogs, prepared for stewing, happened to be situated in a dish, one of them in contact with a scalpel, near an electrical machine, from the prime conductor of which sparks were proceeding. Under these circumstances it was observed, that every spark was followed by convelsive motions in the limbs in the vicinity of the scalpe's The effect was greatest in any limb to which the scaipmight be most approximated. The annexed figure well convey an idea of the manner in which the frogs were situated, relatively to the scalpel; and likewise to the conductor of the machine and knob, between which the sparks were thrown. Galvani was absent, and was indebted for his knowledge of the fact to Madam Galvani, by whom it was noticed.

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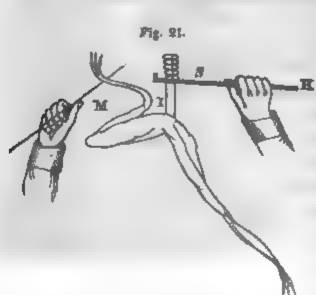
424. The convulsions thus observed, it is now known, were the consequence of the limbs and the scalpel being inductively electrified. It is by an analogous inductive influence, that when one person is struck by lightning, by-standers are stunned, though not in the circuit. Electrical discharges deleterious to human life have been attributed to an inductive charge and discharge consequent to lightning, and analogous to those which convulsed the frogs. (See Dynamic Induction.)

425. Galvani having been previously engaged in the experiments, to verify a conjecture of his, that muscular action is due to electricity, became inspired with additional zeal by these facts, for a knowledge of which he was indebted to accident and his wife's observation. Nevertheless, he would have attributed little importance to the convulsions produced in the frogs, had he not erroneously supposed that the animal organs were instrumental in the generation of the electricity by which they were affected.

426. But, as already stated, (309) the limb of a frog, with the spinal nerves denuded, is a most sensitive galvanoscope, since the minutest discharge of electricity through a circuit of which the nerve forms a part, will produce convulsions in the rest of the limb. Of course during their exposure to the alternate charging and discharging of the machine, the frog limbs acted merely as galvanoscopes.

427. It was, however, subsequently ascertained by Galvani, that convulsions ensue, on making a conducting communication by a metallic wire between the feet of a frog and a portion of the spine appended thereto by the denuded nerves (309). This he correctly ascribed to a power in the parts of the animal to generate electricity;

tween the mode of its previous existence, and that of the charge of a Leyden phial. This view of the subject we not unreasonable at a period when no knowledge had been obtained respecting the electrolytic process, by which the current in a galvanic circuit is generated. But the interfact, that galvanic discharges causing convulsions are producible in three different ways, of which only the last described is due to the animal organs. Now, it is known the leg of a frog may be convulsed, not only as about described, by a metallic communication between the spaninerve and the feet, but likewise by bringing those parainto contact without the intervention of a metal.



lustrate the mode by which the leg of a frog being made to touch the spinal nerve convulsions will ensue a consequence of an inherest diversity as respects electropolarity in the different parts of the animal.

limbs are suspended by means of a glass rod, it. held by the hand at one end.

while the other is situated between the two partially denuded nerves of the spine, a portion of the latter, & which is retained, serving to keep them united. of another glass rod, M, one of the legs is brought at L into contact with a nerve. By these means the limbs may The previous immersion of the be sensibly convulsed. limbs and nerves in brine, is recommended by Sturgeon. 45 rendering success more attainable; but there seems to be some objection to the assistance of an inorganic reagent in an experiment intended to prove that the organs coployed are per se, equivalent to a galvanic triad or elementary battery: it is nevertheless certain that much men active convulsions may be produced by the discharge from the most minute galvanic triad or the smallest spark arising from the friction of an electric (E, 211). Where a prepared frog is made, like the tongue in the experiment of Fultzer, to perform the office of the liquid reagent in a galranic triad such as above mentioned, the slime of the aniral takes the place of the liquid, and the muscles included tre convulsed by a discharge analogous to that by which

he tongue is affected by the appropriate sensation.

430. Resting his inferences upon the superior efficacy extraneous sources of electricity, Volta succeeded in reating an almost universal impression, that the idea of a current originating in the organs of frogs had been erroteous. This impression was strengthened by the, subsequently discovered, surprising energy of his inorganic pile effecting ignition and chemical decomposition.

Of the Diversity in the Sensations produced in the Human Frame by variations in the size and number of the Galvanic Pairs employed.

- 431. The different modes, as respects size and number, in which galvanic apparatus may be diversified, has been alluded to. It was soon found that the shock given to the animal frame was increased more by the number, than by the size of the plates. I have found the shock from two hundred and eighty triads, each two inches square, more severe than from one hundred, nearly eight by fourteen or sixteen times as large. The sensation from the latter is more excruciating, if the contact be sustained, because the quantity is so much more copious; but after repose, the poles of the more numerous but smaller series becomes as much greater than the other as twenty-eight is to ten. This greater intensity enables the charge to act more suddenly, and thus to produce the jerking sensation, usually called a shock, in a much higher degree. Davy mentions that severe shocks were received from his large pile through the floor upon which it rested. I have experienced shocks in like manner, when the floor on which a series of three hundred pairs rested, was wet.
- Dre's Chemical Dictionary, so much of the phenomena of this branch as are associated with experiments on recent carcasses, whether of cold or warm blooded animals, are treated of under the head of galvanism, while the other phenomena are treated and under that of voltaic electricity. It appears to me, that galvanism or galvanic electricity, should be known under the general appellation, and that sufficient justice will be done to the other celebrated inventor, if we speak of the compound circuit, the voltaic series, pile, or battery. The appellation, voltaic series, seems to be preferable, being most descriptive, since whatever may be its construction, it must be massist of a series of elementary batteries.

432. Even with closed eyes an ideal flash of light a produced in the Sultserean experiment. According to Sturgeon, it is best executed when one of the pieces of metal employed is pressed close upon the gums besed the upper lip, the other on the tongue. This preparate being made, a flash is produced by bringing their extensedges into contact. To enhance the effect, the applicate of brine to the tongue is recommended. It should be derstood that the taste endures so long as the contact continues, but the affection of the optic nerve and the such quivering of the tongue are transient effects, attendant to the peculiar intensity of the discharge which takes place

when the circuit is first completed.

433. To a cut sufficiently deep to draw blood, the me feeble galvanic discharge is perceptible; and that of extensive battery is excruciating. On the back of hand, from the comparative delicacy of the epidermis. the superior conducting power of the veins, the sensibility to galvanic discharges is much higher. Moistening the skin, more especially if any saline solution be employed causes such discharges to be much more difficult to best One of the best modes of taking the shock is to comes each electrode with a cup of mercury covered by water Dipping a finger of the right hand in one, and of the in the other, the shock will be felt in perfection. However large the plates of a voltaic series, they cannot promo: any perceptible effect until the number of them, made in operate on a conductor, is sufficient to cause a discharge through it. Hence, they may cause the deflagration of the wire, when they cannot give a shock to an animal ner # compose any chemical compound. When a number or petent to effect a passage is employed, the shock is not w much greater in proportion to the size of the plate b might be expected, just as in passing a needle through a muscle, our suffering would not be much increased. 4though, by doubling the diameter of the instrument. \*\* should quadruple its bulk.

434. The sensation created by a comparatively surfunction of large pairs is more incessant, less jerking, and becomes intolerable from a sensation like the boring of a hot iron. This burning sensation is not surprising, what it is considered that fluids grow hot under the influence of such a series. In 1839, I assisted in some experiments of

the corpse of a recently hanged murderer, of whom the nuscles of the face and those on which respiration is lependent, were made to act. I then observed that ignizion took place whenever the lifting of the end of either lwire acting as an electrode, caused an interruption of the sircuit. The series employed was a deflagrator of three nundred pairs of seven by three.

# Of the Statical Effects of Galvanism.

435. It will be well to remember, that under this designation are placed those phenomena of galvanism which, although extremely feeble as to comparative intensity, still esemble the phenomena of frictional electricity as respects the spark, the separation and approximation of light bo-

lies, and the charging of the Leyden phial.

436. Of all the phenomena of galvanism, only those of a statical kind are to be produced when the circuit is incomplete; and it is under these circumstances, and others which are inconsistent with the other powers, that those in question appear to most advantage. It is only on breaking contact that galvanic apparatus, in the usual forms, will produce a spark. If we attach a metallic mass to one pole of a galvanic battery, and after bringing into contact with it the end of a wire, rod, or strap of metal, in due connexion with the other pole, suspend the contact, a spark ensues; yet none can be perceived on renewing the contact.

A37. Sparks arising from the rupture of the circuit are advantageously seen, by attaching to one of the electrodes a rasp or a rachet wheel, which is so situated as to touch a spring strap in due connexion with the other electrode. By scraping the rasp with the strap, or if a rachet wheel be used, causing it to revolve against the strap, a spark occurs as often as the strap passes from one tooth to another. The spark may be produced by metals, which, like gold and platinum, are difficult to oxidize, but is larger and more brilliant when iron, steel, or mercury, is used. Of these metals, minute portions are mechanically abraded, the combustion of which cooperates in the production of intense light and heat.

438. The phenomenon which, when produced by the puny apparatus of man, we call a spark, is designated as

lightning, when evolved by the gigantic apparatus of m ture. In no respect are the powers of galvanic batters more discordant with those of electrical machines than a respects the spark. By electrical machines of the larger size, sparks of from twenty inches to two feet have been emitted: from a very small one, they may be obtained sometimes two inches in length; yet, before contact the most powerful voltaic batteries could not, by the most genious and skilful efforts of Gassiot, be made to give as apark; and, according to Davy, the battery of the Rott Institution, comprising thirty-two thousand square with of zinc surface insulated by porcelain, did not give a spea at a greater distance than one-fortieth of an inch (36%) It follows, that to give a spark at ten inches, or four dred times as far, would require a striking power with would be to that of Davy's series as the square of for hundred to that of one; so that the striking power of t machine giving such a spark must be one hundred sixty thousand times as great as that of the celebrate battery above mentioned.

439. With insulated voltaic batteries of 1626 pairs made by Crosse, and of 3520 pairs made by Gassiot, excited in river water, sparks were obtained before contact. From the last mentioned battery we are informed that a correspond to them was obtained. The other powers of the batter are enfeebled, in proportion as its statical power is

strengthened.

the dry pile of De Luc, as modified by Zambon. According to Daniell, when a wire proceeding from one proof a pile of 20,000 pairs, was drawn lightly over a mished metallic surface communicating with the other properties of minute sparks were obtained. A Leyden we tery might be so charged as to produce a disagreeable shock extending to the elbows and shoulders (375).

441. It does not appear to me that sufficient consideration has been given to this form of voltaic apparatus by the advocates of the purely chemical origin of the galvatus of rent. Faraday, the, matter of fact, Ajax of that corps is sists on the necessity of a species of electrolytic convects between the zinc and copper surfaces, to the conveyance of the current. He urges the necessity of a process had that which is supposed to take place during the decomposition.

pretended, that any such electrolytic series of decompositions can go on by means of the minute and detached particles of hygrometric water contained in the paper material of Zamboni's disks: and admitting that moisture, thus existing, could, by its electrolytic decomposition, be adequate to sustain the electrical discharges for years (381), by what process is it transferred from surface to surface through the intervening paper? Evidently it cannot be conceived to pass by the series of decompositions and recompositions in rows of the atoms of an intervening liquid. Sturgeon avers, that the powers of De Luc's series is greater in proportion as the surfaces are kept brighter, so that chemical action, far from being necessary, is actually injurious to the energy of the apparatus.

# Of Continuous Ignition.

442. I deem it expedient to distinguish, under the general designation of continuous ignition, those cases in which a body, to be ignited, is a good conductor (a wire or charcoal, for instance), forming that part of the galvanic circuit which lies between the electrodes (356). In this case a wire being uniform in diameter and shape, the length may be directly as the intensity. The larger the plates, the stouter the wire which can be rendered incandescent: the greater the number of the series, the longer. Within any reasonable limits, the sectional area for a given length may be as the area of the oxidizable surface of one pair; the length as the number of pairs in the series.

the length as the number of pairs in the series.

443. By the gigantic apparatus of Children, a platinum wire, five feet six inches in length, and more than one-tenth of an inch in diameter, was heated red-hot. By my deflagrator, several feet of wire of the same material may be raised to incandescence, and when consisting of iron, it may be burned by the atmospheric oxygen before fu-

sion.

444. The platinum wire employed in igniting gaseous mixtures (C, 335), is liable to fusion, if care be not taken to break the circuit before the temperature reaches its maximum.

445. It is remarkable that if the wire, subjected to continuous ignition, be of uniform thickness, as is usually the

case with a wire in good preservation, the ignition began about the middle, and extends gradually towards the dotrodes.\*

connect the poles of a powerful deflagrator, on competate the circuit, by bringing the plates and acid into counted the charcoal will be instantaneously ignited throughout

# Of the Ignition of a Continuous Conductor by a Calorinate or Deflagrator.

447. If the electrodes of a galvanic triad, of the construction to which the name calorimotor has been guestic be united by a wire before the immersion of the plates; or immersing them, the wire will be more or less heated, and the size and length of the oxidizable metallic surface, and the size and length of the wire; excepting that the netensity produced by one pair lessens as its size is enlarged. Although on estimating the heat which it can produce, a wire of a certain magnitude, it may be found in quantity proportionable to the size, in order that this proportionable quantity may appear, the wire must have magnitude to the size of the size o

As, when the wire is of a sufficient length, it takes some time for the whole become incandescent, and as fusion cannot be effected in the central portion as leg as the rest of the wire is below a white heat, the operator is enabled to avoid from the central portion, while raising it, nevertheless, to the highest temperature what platinum will bear. To effect the combustion of hydrogen with nate is not a quires a heat so intense, that I always use a wire for that purpose of near its inches in length, and of about No 26 of the wire gauge

An argument in favour of the theory of Dufay has been founded upon the tenthat the heat taking place midway corresponds with the meeting of two fluids it is intermediate space within the circuit equidistant from their respective sources. But should be kept in mind, that agreeably to the theory of two fluids the negular end of the pile has to receive as much vitreous as it yields up realn us and the put two end has to receive as much resinous as it emits of vitreous electricity.

It follows, that the conditions necessary to electric activity, agreeably to the theory, consist in the simultaneous motion of two fluids in opposite directors. It cannot consist in their coming together at any particular part of the electrons than in any other. It is true, that when the circuit is interrupted, an excess vitreous electricity will be accumulated towards the positive end of the series an excess of the resinous towards the negative end, but, on completing the commensurate with the discharge of the vitreous accumulation into the action be a discharge of the resinous electricity in the opposite direction. In the commensurate with the discharge of the resinous into the cathode with the discharge of the resinous and the cathode with the upper the circuit, to restore the equilibrium which the electro chemical reaction is necessarily and the reaction is necessarily destroying

The phenomenon under consideration, though irreconcilable either with the there of one or that of two fluids, agrees with the idea of waves of polarization moting from the centre of the generating battery towards the extremities, thence passage through the metallic conductor to neutralize each other at an intermediate point the wire. Possibly each wave, carrying with it calorie in a latent state, may risk it up when, by neutralization, the power to retain it ceases; and thus may arise the heat evolved at the place of meeting, so long as the current is sustained

great to be consistent with intensity. Hence, a more intense heat may be produced in a fine wire by a triad of a square foot of oxidizable surface than by one of fifty. On this account it is necessary in a very large calorimotor to employ more than one triad in order to produce a white heat in a wire whether stout or fine\* (331).

# Of Deflagration.

448. Those experiments in which the contact or continuity of the conductor between the electrodes, after being established, is destroyed by its reaction with the current, or by the interference of the operator, fall under the head

of deflagration.

449. Thus when suitable tapering pieces of charcoal, severally attached to the electrodes of an active and sufficiently powerful series, after being brought into contact, are separated, the resulting interval becomes occupied by an arc of flame of which the light and heat are the most intense that art can produce. It is not without pain, and the risk of permanent injury, that this flaming arch, or

For more than twenty years, I have employed calorimotors for effecting the explosive combustion of gaseous mixtures, or for igniting a jet of hydrogen in oxygen in the Lavoisierian experiment for the synthesis of that liquid (834-5). In such experiments, that distinguished philosopher, and generally all the chemists of the old world, have used for the same object, the electric spark. This it is difficult to command in damp weather, and in the vicinity of the water usually employed in pneumato-chemical experiments.

In June, 1831, I was induced to make a series of experiments, in order to explode the charges of gunpowder in rock-blasting, and succeeded by one of my deflagrators in exploding twelve charges at the distance of one hundred and fifty feet. Of these experiments, I published an account in 1833, in the Franklin Journal, and recommended this process as peculiarly well suited for blowing up fortifications and blast-

ing under water.

In England, about five years afterwards, efforts having been unsuccessfully made by other means of ignition, to remove the wrecks of vessels by the explosion of gunpowder previously introduced into their holds, galvanic ignition was resorted to pursuant to my suggestion. Nevertheless, upon the plea that my apparatus had not been employed to affect ignition "under water," resort was had to another, which was of course, previously to trial, amenable to the same objection. In fact, the deflagrator was peculiarly competent to effect ignition under water, far more so than the constant battery which was used. This arises from the completion of the circuit being affected by throwing the acid on the plates at the moment when the ignition is required. In this mode of operating, a transient power may be obtained far greater than can be had from an equal extent of surface in a constant battery.

Finally, however, Col. Paisley succeeded in blowing up the wreck of the Royal George, by means of galvanic ignition. Since then, a person by the name of Colt, has applied the process of galvanic ignition so successfully, as to blow up a versel at the distance of several miles from the battery employed. In the employment of this process, it does not appear, that he made any due acknowledgment either as to Col. Paisley or myself. So entirely was the committee of congress with whom he conferred, in the dark, that in April last, both Prof. Joseph Henry, of Princeton, and myself, were applied to, by the Secretary of War of the United States, to give that information, which it was evidently Colt's duty to have given at the outset.

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450. On comparing the light thus generated from particles hundred pairs of seven by three inches, with that of access dle, the ratio appeared to be as sixteen hundred to a The shadow of the whiskers, or of a prominent loss of hair, partially intercepting the light, was cast upon a wal with extraordinary accuracy. The side of an adjaces house, on which the rays fell in the night, appeared as i

illuminated by the sun-

451. When a rod of platinum, of a quarter of an inc in diameter, while attached to one pile, was made to took charcoal attached to the other, the instantaneous fusion of the platinum at the point ensued. A knitting needle sestituted for the rod of platinum was explosively defegrated. In such cases the wire is destroyed in detail, the combustion being limited to the vicinity of the end. For this and other deflagrations where a metal is in questian it answers better to have one charcoal electrode, than w have both of metal. Two metallic electrodes are hable w be soldered or welded together. The incorporation the arising, takes place even under water. A platinum wat brought into contact with an iron rod under water may be welded thereto. It is preferable to have one wire always stouter than the other. Of course tin-foil and other left metal are rapidly deflagrated by the same process as the knitting needle. A jet of mercury, so regulated as not be form a stream quite continuous, being made the medium of the discharge of a deflagrator, a splendid effulgence of sues, arising from the combined effects of the combustice of the metal and the galvanic light.

# Deflagration with Liquids.

452. Let a concentrated aqueous solution of chloride of calcium be first brought fully into contact with a stoat platinum wire, acting as the positive pole of a deflagrates

deterred from subjecting his eyes injudiciously to an ordeal like that which may

had undergone

<sup>\*</sup> On one occasion, after making some experiments with an extensive series at eyes were so much affected, that all other flame appeared of a deep to see that on the following day they were blood shot, and a preatly inflamed, that a we deemed expedient to apply twenty leaches to each to procure relief.

On another occasion, similar consequences ensued to an assistant, who was >=

f about three hundred pairs, seven inches by three, in full peration. In the next place, let a platinum wire of about 10. 30 of the wire gage, acting as the negative pole of the ame series, be allowed barely to touch the solution, the nost intense ignition will ensue, causing the platinum to e fused rapidly into globules, which will be found at the ottom of the solution. An analogous result ensues with ther saline solutions, if capable of a similar degree of ensity. The fusion of the platinum, however, is much nore readily effected with the chloride of calcium. This rises, I infer, from the extrication of the calcium, its union rith the platinum momentarily, and subsequent rapid commation by the oxygen of the water.

453. It is remarkable that this experiment could not be erformed by a deflagrator of one hundred pairs of seven nd a half by fourteen, although succeeding perfectly with three hundred pairs of seven by three, having less han two-thirds of the surface disposed in a series of an

nferior number.

454. There is no mode by which a small portion of a ubstance can be subjected to heat in the absence of the xidizing influence of the air, or with the co-operation of he deoxidizing affinity of hydrogen, so powerfully, as that of a large voltaic series acting in a receiver capable of \*xhaustion and replenishment with that gas or any other. Subjecting phosphuret of lime in this way to the intense gnition produced by two hundred pairs, each of nearly even and a half by fourteen, I reduced the calcium very nearly to a state of purity, as a small portion acquired a metallic brilliancy under the burnisher, and decomposed water rapidly on being thrown into it.

455. Having, in like manner, exposed cyanide of calcium, the result was similar, but less exceptionable, as this compound does not possess the property of liberating hydrogen from water, so that the acquisition of the property indicated more certainly that the substance evolved was calcium. In either case lime was found in the water, in which the supposed calcium had been immersed.

# Of the Electrolytic Power of Galvanism, or Galvanic Power of Chemical Decomposition.

456. The voltaic series was first discovered to have powers of decomposition, or electrolysis, by Nichola and Carlyle. These ingenious investigators found a subjecting water to copper wires acting as the electres of a voltaic pile, that while hydrogen was evolved at the negative electrode, the positive electrode was oxident But when wires of gold or platinum were used, both of the gaseous elements of water were evolved. In the follows: summer I was invited by my predecessor, Dr. Woodbotte to see the experiment then going on in his laborators. means of a series constructed of disks of copper, zuc. 131 moistened cloth. This was considered as a great step ! science at that time, when it was still denied that with could be decomposed, or that the hydrogen obtained ! passing it in the state of steam over incandescent was an element of that liquid. To the evolution of the gaseous elements by the voltaic pile, those who considered it as a compound, appealed with elation; and the proces that it could be thus electrolysed, as we would now sywere finally such as to silence all who had been increalous.\*

Having to illustrate the decomposition and recomposition of water to a come between three and four hundred pupils, I have found it expedient to exhibit the recess on an extensive scale. Hence, for many years, I have employed a grant of about two inches in bore, and about two feet in height, furnished with the lures, B, b, about three inches below the upper extremity, where the bore country to an apex, having an aperture not larger than a goose quilt. I por this apex the is an iron cap, in which a female screw is wrought so as to allow a large than cock, C, to be screwed into it

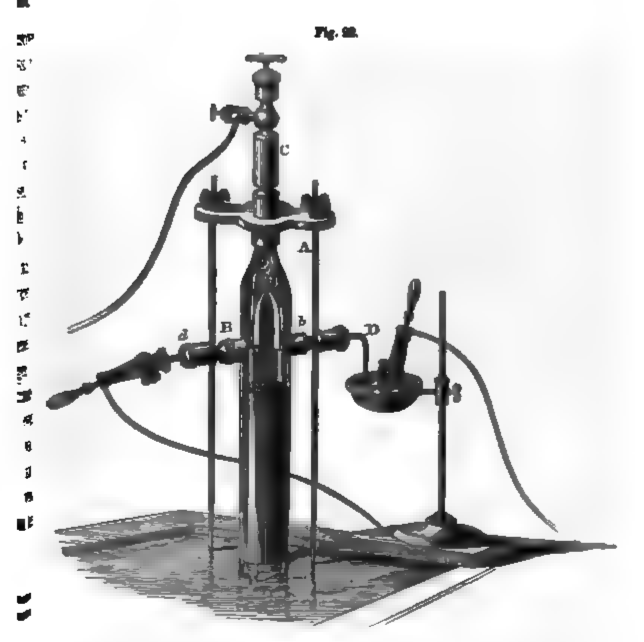
Upon the tubulures from caps are comented, which are so wrought, as a time aid of appropriate screws, to constitute stuffing boxes. Through each of translations rod, D, or d, is introduced, and fastened to plates of platina, to act as trodes," agreeably to the language of the celebrated Faraday.

The tube being supported over the mercurial catern, by means of a commentation with an air pump through the valve cock and flexible leaden pipe, the box of the tube is a substitute of air so as to cause the mercury to take its place. The state of the place.

The tube being supported over the mercurial c stern, by means of a constitution with an air pump through the valvo cock and flexible leaden pipe, the best the tube is exhausted of air, so as to cause the mercury to take its place. The secury is so far displaced by a solution of borax, consisting of equal parts of water is a saturated solution of that salt, as to sink the surface of the column of metal 2 to tube about an inch or more below the "electrodes". The projecting end of securities rods, D. d. to the other ends of which the "electrodes" are severally attacked is bent at right angles outside of the tube, so as to enter some mercury in an iron are sule, supported purposely at a proper height, and communicating with one cut my deflagrator of an hundred pairs of Cruckshank plates, described page 22. The ourse the rod of the other electrods must have a communication with the other seconds.

<sup>\*</sup> Engraving and Description of an Apparatus for the Decomposition and Recomment of the Laboratory of the Medical Department of the two sity of Pennsylvania.

457. In 1807, Davy made the grand discovery of the com-"position of potash and soda, by means of two hundred pairs



of the deflagrator. Under these circumstances, if the circuit be completed by throwing the acid on the plates of the deflagrator, a most rapid evolution of hydrogen and exygen will ensue in consequence of the decomposition of the water, so that within a few seconds, several cubic inches of gas will be collected.

The action being now suspended by throwing the acid off the plates, and the foam

The action being now suspended by throwing the acid off the plates, and the foam being allowed to subside, the resulting gaseous mixture may be ignited, and of course condensed, by completing the circuit again as at first, and at the same time causing the ends of the "electrodes" to come into contact with each other, and thus to produce a spark. This contact is effected by causing a very slight movement in the rod, bent at right angles, and entering the mercury in the iron capsule. Of course the process may be repeated as often as can be reasonably desired.

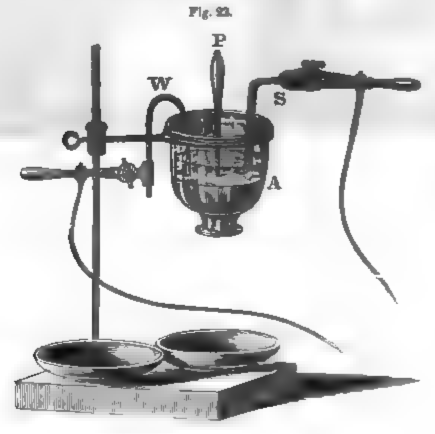
Generation of the Amalgam of Ammonium, upon a large scale, by " Electrolization."

An open necked bell, A, inverted and supported as in the figure, is furnished with a plug, P, by which the crifice in the neck may be closed or opened, more or less, as desirable. The lower part of the cavity of the bell is occupied by mercury, covered by a stratum of a concentrated solution of chlorida of ammonium (sal ammoniae). An iron wire, W, so covered with gum elastic, as to pass through the solution of chloride without any conducting contact, establishes a communication between the secretary and the negative pair of the series. By means of a platina strap, S, a like communication is effected between the positive pole and the dissolved chloride. The circuit being completed, the mercury soon swells up, assuming the well known ap-

ø

of four inch square plates. By Sweiger and Ben analogous results were obtained by using mercury at negative electrode. By the presence of this metal nascent metal of the earth subjected to the process, we a certain degree protected from oxidation, which owise would have prevented it from being retained metallic state. Davy, by submitting the amalgams to tillation, obtained minute portions of barium and a tium, imperfectly deprived of mercury. In the distilt of the amalgam of calcium, he failed, as the tube to before the process was complete. From my experimental infer, that with sixty grains of mercury, which was the could not have obtained available portion of the metal had the process success.

458. By means of two deflagrators, each of one but pairs, seven and a half inches by fourteen (375), I enabled, in 1839, to procure as much as 3600 gmi the amalgam of calcium. From this the mercury was



pearance of an amalgam. When a sufficient quantity of this is formed by ico the plug, any excess of mercury may be allowed to run out into one receptac afterwards the amalgam may be received in another, care being taken tick aperture, by means of the plug, before the residual aqueous solution can took these means about half a pint of the amalgam has been obtained. The disserted bell, which I have used, is about five inches.

It has been found advantageous to add liquid aminonia to the solution is cient quantity to prevent the evolution of free acid, which seems to destroy the gam when in contact with it beyond the influence of the circuit.

tilled, by means of a crucible included in a cast iron ralembic, the cavity of the latter having been previously supplied with some pure caoutchoucine to expel oxygen. In this way I obtained sixty grains of calcium. In subsequent efforts I also succeeded in obtaining about the same quantities, respectively, of barium and strontium.\*

# On Secondary or Indirect Decomposition.

459. In the case of the decomposition of a liquid by wires, acting as electrodes, it was formerly inferred that the elements which appeared at the electrodes were liberated by a direct electrolytic influence, but Faraday has shown that the evolution of the element of one compound, another compound being present, may cause the decomposition of

The cuts having unexpectedly been found to be missing, and not deeming it expedient to stop the press in order to replace them, I have to omit giving an engraving and description of the apparatus employed in the processes above mentioned (444, 458).

As no account of these metals, as procured by me, is given under Inorganic Chemistry, the statements made respecting them in the last edition of this Treatise are

subjoined.

#### Properties of Calcium, Barium, and Strontium.

When the heat was sufficient to expel all the mercury, the metal was found adhering to the bottom of the crucible in a crust, which required an edge tool to detach it, though no incorporation of the iron with it appeared to have taken place. When in distilling calcium, a crucible of platina was employed, a portion of this metal was found to have united with some of the calcium, being detached therewith in the form

of a bright metallic scale.

In consequence of their susceptibility of oxidation, and of union with the elements of naphtha, the metals obtained as abovementioned were devoid of metallic lustre until their surfaces were removed by a file or burnisher. Either was rapidly oxidized in water, or in any liquid containing it; and afterwards, with tests, gave the appropriate proofs of its presence. They all sank in sulphuric acid; were all brittle, and fixed, and for fusion, required at least a good red heat. After being kept in naphtha, their effervescence with water is, on the first immersion, much less active. Under such circumstances, they react at first more vivaciously with pure ether than with water, or even chlorohydric acid; because in these liquids a resinous covering, derived from the naphtha, is not soluble, while to the ether it yields readily.

By means of solid carbonic acid, obtained by Mitchell's modification of Thilorier's process, I froze an ounce measure of the amalgam of calcium, hoping to effect a partial mechanical separation of the mercury, by straining through leather, as in the case of other amalgams. The result, however, did not justify my hopes, as both metals were expelled through the pores of the leather simultaneously, the calcium forming forthwith a pulverulent oxide, intermingled with and discoloured by mercury in a

state of extreme division.

By the same means I froze a mass of the amalgam of ammonium, as large as the palm of my hand, so as to be quite hard, tenacious, and brittle. The mass floated upon the mercury of my mercurial pneumatic cistern, and gradually liquefied, while

its volatile ingredients escaped.

When the freezing of the amalgam was expedited by the addition of pure ether, the resulting solid effervesced in water, evolving ethereal fumes. This seems to show that a portion of this ether may be incorporated with ammonium and mercury, without depriving the aggregate thus formed of the characteristics of a metallic alloy.

nant affinity for one of the constituents of the composite to which it is thus presented. Thus the deposition of metal at the negative electrode from a solution of metal at the negative electrode from a solution of metal its salts, by exposure to the circuit, may be the constituence of its oxide being decomposed by the affinity of hydrogen in a nascent state as evolved from water. The nitrogen evolved at the anode from liquid ammonia, most like circumstances, is conceived to be liberated by the union of its hydrogen with the oxygen of water dream.

decomposed.

alkaline earths, when mercury is employed as the next tive electrode, an amalgam is obtained of the metalic radical of the salt, I had formed the opinion that t is the metallic oxide which is the subject of direct decomposition, and that when hydrogen appears at the control of the thode it is the consequence of the decomposition of the water by the metal. It is known that as soon as either of these in its metallic state touches water, seizing the oxygen, it liberates the hydrogen of this liquid. Subsequently, I found that Matteucci, a distinguished Italian pholosopher, had arrived at the same opinion as myself, of this subject, after performing a series of laborious experiments with a view to ascertain the truth.

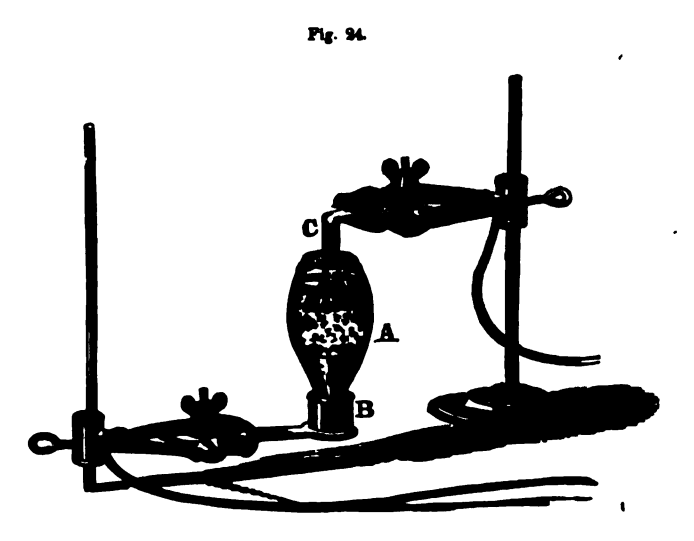
461. This question must excite great interest from 's fact, that it has an important bearing on the electro's investigations recently made by Daniell, which by Kasa Graham, and others, as well as the author of them, have been considered as justifying the salt radical theory. But in either way of viewing the subject, Faraday's suggestions that the elements, which are the ostensible products of the process, may be indirectly evolved, is fully sustained.\*

<sup>\*</sup> Figure 24 represents an apparatus which may be conveniently employed for a electrolytic formation of amalgams with the metals of the alkalies or and reserves.

The lower end B of the receiver A is perferated so as to have an iron error ferably a-platinum wire, so luted therein, that one end may just enter the car'r' the receiver, while the other end is soldered to a metallic strap supporting the ceiver, and held by a vice communicating through a wire rope with the positives of a competent voltaic series. The lower part of the cavity of the receiver is cupied by some mercury, necessarily in contact with the end of the wire alrest described as entering that space. Upon the mercury a stratum of the aqueous sometion containing the desired metal must be situated. By means of a strip of plates in contact with this solution at one end and at the other held by a vice in communication with the positive end of the voltaic series, the circuit is completed. For

# ults obtained by long enduring Galvano-Electric Currents of Low Intensity.

162. Becquerel and Were Fox, have drawn the atten1 of the scientific world to processes, in which galvanic
1 rents long continued, of low intensity, have been suc1 study employed. It has been rendered probable by
1 investigations and observations of Were Fox, that gal1 ic circuits are among the most efficient causes of
1 tallic depositions in mines, and by Becquerel, the pro1 ses of nature in the formation of crystals have been
1 tated. In this path, laborious experiments, of unpre1 ented endurance, have also been made by Crosse. It
2 during some efforts to produce by the voltaic current
1 stals, and other compounds, analogous to those found
1 nature, that this zealous electrician observed a species
1 louse, since called acarus galvanicus, to appear. It is
1 tainly miraculous that these insects should come into
1 stence under the influence of agents, which are never
1 nd to enter into the constitution of animals, and which
1 of a nature to destroy life in the case of animated



s circumstances, the series being in a sufficient degree of activity, the mercury ig as the negative electrode separates, and unites with any metallic cathion in electrolyte held by the solution subjected to the process; meanwhile the anion se electrolyte is evolved at the anode, or where the surface of the platina electron is in contact with the liquid (358).

nature in general. The insect so much resemble to cheese mite, that it might be mistaken therefor out

hasty examination.

with modifications, by Weekes of the London Electral Society. This ardent investigator subjected to the advance current solutions of silicate of potash, in glass receivers over mercury, the greatest care being taken guard against the access of any extraneous bodies. It der these circumstances, after an uninterrupted exposm during upwards of a year, insects are alleged to have make their appearance perfectly resembling those which seemed to come into existence during the experiments of Cross. Mr. Weekes has since asserted that the acari appeared to have originated in a solution of ferroprussiate of potast. The origin of these insects is inexplicable, since it is utterly impossible that they can be created by the agent and process employed.\*

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\* On this subject, in my introductory, delivered in 1843, I expressed system follows:

A few years since, many of the more inconsiderate among the reading point of society, were carried away by an account respecting certain experimental therestions made by an enthusiastic electrician, to believe in the generation of certain sects through the influence of the voltaic current. It struck me at first, that a nothing less than the mirroulous interposition of the Deity could give rise to and life, this narrative must be a jocular fiction, like the statement respecting countries, this narrative must be a jocular fiction, like the statement respecting countries, this narrative must be a jocular fiction, like the statement respecting countries, this narrative must be a jocular fiction, like the statement respecting countries, this narrative must be a jocular fiction, like the statement respecting countries, this narrative must be a jocular fiction, like the statement respecting countries.

It was, nevertheless, true, that an equivocal description had been given against earnest, tending to create the idea that animals had been generated by a vive or rent out of salicious earth, alkali and acid—It subsequently appeared in that there species of start or lice had been created, but that they had been stown ates as the themselves by the agents and apparatus employed. According to my appreciable it had been quite as practicable to repeat the Mosa commacle of thinging start we a rock, by the blow of a rod, without supernatural and as to create sent enterprise beings by the materials and apparatus employed by the electric and Farm we conable were it to expect the formation of a with by a specting the oresis metals requisite for its construction to the voltage process.

Perhaps turns n gentus has an no instance approached more nearly to be prestion of a machine beging the attributes of an armin than in the construct a 1 at locometive steam engine, of which the furnace may be considered as performed office of a stomach, the cyander and pipes that of the bowels, wante the start conand wheels act as limbs—but still the will the instruct, the power of scalespear

tion, and the heaven born sense of existence, are deficient.

In fact, human efforts have never attained the point of making a self-miral machine, or perpetual motion, however wrong-headed projectors may have serious themselves and others with false expectations of success in that impracticable per ject. For, although there he in nature incessant movements, like those of the planets, or of waves, &c., and currents which, originating from calorific, or electrochanges, move or flow without intermission from age to age, such sources of movement to be created by man

## Important Inferences of Faraday, respecting Electrolysis.

464. Among many important services rendered by Faraday, is that of temonstrating that electro-chemical decomposition takes place in equivalent atio, being always directly as the quantity of electricity which passes by he electrolytic separation and rearrangement of the ions (360). For the avestigation of the phenomena dependent on this law, an instrument was levised by him, which, being interposed in the circuit, enabled him accumtely to measure the gaseous products of the electrolytic decomposition of vater. The result, thus obtained, was assumed as a standard by which to stimate the products of the decomposition of other electrolytes exposed to he same current simultaneously with water. To the instrument thus employed the name of volta electrometer, or by abbreviation voltameter, has seen given. The construction of this instrument does not differ essentially from the apparatus above described for the decomposition of water (note to 150), excepting the absence in the latter of graduation, by which the volume of the gas generated is ascertainable on inspection.

465. By means of this apparatus it was ascertained, that when the fused hlorides of tin and lead were either severally or simultaneously subjected to the same active circuit, so that the products of their decomposition might compared with those of water with water, the weight of the elements, wolved from this liquid, collected in the voltameter, and those of the chloine and metals, liberated from the chlorides, were to each other as their

quivalents, H 1, O 8, Cl 36, Pb 104, Sn 59, C (560).

466. By these results, and many others of an analogous character, it seems to have been fully established, that the quantity of any electrolyte rhich can be decomposed by a galvanic current is directly as the quantity of the electricity which passes through it: moreover, that the weights of different electrolytes discomposed by the same quantity, will be to each other, a the numbers by which their atoms are represented in the table of equivalents. The only motive for increasing the intensity by a numerical extension of the series employed, is that for the passage of the fluid by the electrolytic process, the requisite force must be as the reciprocal affinity of the one of the electrolyte employed. Thus, as in the chloride of calcium, a tronger affinity is to be overcome than in decomposing the iodide of that notal, a greater number of pairs are required, but the quantity of the compounds in question decomposed, will be, ceteris paribus, as the size of the plates, without reference to the number of the series.

467. Not only did Faraday show that the weights of the ions evolved by the same galvanic current from different electrolytes, and consequently he weights of these, were to each other as their equivalent numbers, it was also shown by this skilful experimenter, that to produce the same quantity of electricity from different metals by galvanic reaction, the weights of the quantities corroded, must be to each other as the equivalent numbers of the

netala.\*

It seems to me remarkable, that under these circumstances, no reference is made by Faraday to the analogous results, obtained by Pettit and Dulong, respecting specific seat. It is well known, that it was rendered highly probable, by these philosophers, but the quantity of heat given out by bodies was equally great for each atomic squivalent. The specific heat of every gas is the same, so that although the weight a equal volumes of gaseous hydrogen and oxygen are as one to sixteen, the quantity

#### An Interesting Summary of Faraday's Inferences.

The following summary of his inferences is given by Paraday.

488. "i. A single ion, i. e. one not in combination with another, with have no tendency to pass to either of the electrodes, and will be painty indifferent to the passing current, unless it be itself a compound a malementary ions, and so subject to actual decomposition. I put the next founded much of the proof adduced in favour of the new theory of combined decomposition, which I put forth in a former series of the searches.

469. "ii. If one ion be combined in right proportions with cathestrongly opposed to it in its ordinary chemical relations, i. e. if an ent be combined with a cation, then both will travel, the one to the cath, to other to the cathode, of the decomposing body.

470. "iii. If, therefore, an ion pass towards one of the electrode, a other ion must also be passing simultaneously to the other electrode, a though, from secondary action, it may not make its appearance.

471. "iv. A body decomposable directly by the electric current, a carellectrolyte, must consist of two ions, and must also render them up des

the act of decomposition.

472. "v. There is but one electrolyte composed of the same two dimentary ions; at least such appears to be the fact, dependent upon a little that only single electro-chemical equivalents of elementary ions can get

the electrodes, and not multiples.

473. "vi. A body not decomposable when alone, as horacic acd, we directly decomposable by the electric current when in combination. it makes as an ion going wholly to the anode or cathode, but does not write its elements, except occasionally by a secondary action. Perhaps a superfluous for me to point out that this proposition has no relation to a cases as that of water, which, by the presence of other bodies, is realised better conductor of electricity, and therefore is more freely decomposition.

474. "vii. The nature of the substance of which the electrode is provided it be a conductor, causes no difference in the electro-decount tion, either in kind or degree: but it seriously influences, by secondary it tion, the state in which the ions finally appear. Advantage may be use of this principle in combining and collecting such ions as, if evolved a the

free state, would be unmanageable.\*

475. "viii. A substance which, being used as the electrode, can combain with the ion evolved against it, is also, I believe, an ion, and combains such cases, in the quantity represented by its electro-chemical equivalent All the experiments I have made agree with this view: and it seems to at present, to result as a necessary consequence. Whether, in the sectedary actions that take place, where the ion acts, not upon the matter of the

of heat in the equal spaces which they occupy are equal; so that the specific head these gases are inversely as their densities or weights. The densities of hydrogen oxygen, nitrogen, and chlorine, being respectively as 1, 3, 14, 36, the specific head of equal weights of hydrogen, oxygen, nitrogen, and chlorine, are inversely as them numbers.

"It will often happen that the electrodes used may be of such a nature is, which they are immersed, to produce an electric current, either accessing with or opposing that of the voltaic arrangement used, and in this war, which direct chemical action, may sadly disturb the results. Still, in the midst of these confusing effects, the electric current, which actually passes in any aircomption the body suffering decomposition, will produce its own definite electrorise action.

\_=lectrode, but on that which is around it in the liquid, the same consequence

Tollows, will require more extended investigation to determine.

476. "ix. Compound ions are not necessarily composed of electrochemical equivalents of simple ions. For instance, sulphuric acid, boracic excid, phosphoric acid, are ions, but not electrolytes, i. e. not composed of **Blectro-**chemical equivalents of simple ions.

- 477. "x. Electro-chemical equivalents are always consistent; i. e. the remember which represents the equivalent of a substance A, when it is separating from a substance B, will also represent A when separating from a third substance C. Thus, 8 is the electro-chemical equivalent of oxygen, whether separating from hydrogen, or tin, or lead; and 103.5 is the electro-**Ehemical** equivalent of lead, whether separating from oxygen, or chlorine, or jodine.
- 478. "xi. Electro-chemical equivalents coincide, and are the same, with ardinary chemical equivalents."
- 479. It is much to the honour of Faraday that these opinions, which were anostly original, appear to be for the most part sanctioned by the scientific world. The greatest error is, as I conceive, in his language being of a nature to convey the idea of a loco-motion, which cannot take place agreeably to his own premises, and which I infer it was not his intention to treat ne necessary to the electrolytic process. On this, I have already commented (397). Subjoined are some additional strictures (490, &c.).

#### On Sir H. Davy's Experiments on the Transfer of Acids through Alkalies, and of Alkalies through Acids by the Voltaic Currents.

480. When three vessels A, B, C, were associated by moistened strips asbestos, so that the intermediate vessel A, held either an acid or an alkali, B, a saline solution, C, pure water, Sir H. Davy found that by mking them into a voltaic circuit, either an acid or alkali would reach the water of C. Hence it was inferred that, under the magic escorting influzace of the current, an acid ion, as we now call it, might pass through an alkali, or an alkaline ion through an acid. Faraday treats this miraculous result as the natural consequence of the interposition of a body capable of combining with the ion during its journey. It seems, however, inconsistent with his adoption of the Grotherian hypothesis (389), respecting the mode m which ions are liberated by the current, that an ion should actually ravel. Consistently with that view of the subject, in order that an acid ion should be liberated at the anode, and an alkaline ion at the cathode, there must be a row of compound electrolytic atoms, subsisting between the elecrodes, in which the ions evolved are ingredients.

481. I trust that the following quotation from the work of his friend,

Daniell, fully justifies my conception of the subject.

482. "We have now to consider how the impulse, which is derived from the zinc, is transmitted through the liquid to the platinum; and how we may account for the extraordinary fact, that the radical of a portion of the acid combines with the former, while the equivalent hydrogen, with which it was associated, is evolved upon the distant surface of the latter.

483. "No visible transfer of the disunited elements takes place, and if the vessel which contains the acid, be divided either by a diaphragm of bladder, a partition of porous earthenware, or other substance capable of imbibing the liquid, and the two plates be placed on its opposite sides, no impediment arises to this extraordinary separation.

484. "The force must be conceived to travel in this part of its circuit by

us in forming a just notion. When a number of ivory balls are from a pended in a row so as just to touch one another, if an impulse he can one of the extreme ones, by striking it with a hard substance, the fer is be communicated from ball to ball without disturbing them, to the most distant, which will fly off under its full influence. Such as are but remote, and must not be strained too far; but thus we may remote that the force of affinity receives an impulse in a certain direction and enables the hydrogen of the first particle of aqueo-acid which we decomposition, to combine momentarily with the radical of the next per in succession; the hydrogen of this again with the radical of the next second on, till the last particle of hydrogen communicates the impulse in

platinum, and escapes in its own elastic form."

485. In order to render this resort to balls consistent with the phrases. which it is intended to enlighten, I conceive, that instead of supposes row of balls, we must suppose two rows, one parallel to the other. same time, it may be expedient to imagine that one row, represents 2 anions, are white, while the other, representing the cathions, are but and that they are arranged in pairs, each of which represents a concutatom of the electrolyte. It seems necessary, also, that there should a me impulses given, opposite in tendency, so that one operating upon act . may cause the expulsion of an anion, at one end, from the whre real cathion, at the other end, from the black row; the residue rearrant themselves within the same boundaries, so as still to exist in para. In the have no unpaired ball at either extremity. Meanwhile it is inevented 2 this loss of an anion and a cathion will be compensated by the accest another pair, of which, to make the analogy hold, there should be as in dance at hand. Moreover, the row, thus refitted, should, under the 🖘 latory influence of the heat arising from the process, give place to asrow, similarly constituted, and equally susceptible of electrolyzation. The it follows, from the illustration to which Daniell has reserved, when we carried out, that, consistently with his own explanation of the town electrolyzation, there can be no traveiling of an ion from one decree the other. It is alleged by him, that the force travels, per se, by a second convection, without disturbing any other ball excepting that which many at the extremity of the row (484). Hence, in conformity with the terror it seems to me that an acid can only reach the cathode, through no analyses solution, by the endosmosis of the salt and its subsequent decomposition

#### Of the Electrotype or Galranotype Process.

exchange of partners throughout a row of atoms of water extend to one electrode to the other, an atom of oxygen may be literated to anode, and an atom of hydrogen at the cathode (35%). Consistent to a galvanic circuit comprises two electrolytes, separated by a pressiphragm having a common anion, oxygen, for instance, but lift at thious, as, for example, hydrogen and copper, this metal may a traction of the diaphragm perform the office which hydrogen performs on the side. Hence, while at the anode oxygen will still be evolved, a document of copper will, at the cathode, be substituted for the evolution of hydrogen oxide be substituted, the deposition of the metallic radical will take the could of hydrogen, unless the radical be one of those which in

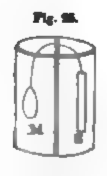
se water on contact, at ordinary temperatures. If it be one of those, by e consequent decomposition of water, hydrogen will appear, as if water are the sole electrolyte. Analogous results will ensue, when any other mple electro-negative element is the common ingredient in electrolytes bjected to the galvanic current while separated by a membrane. Hence, the electrotype process, which it is now the object to explain, only those stale can be employed, of which the deposition in the metallic state is ncticable, in consequence of their not uniting with the exygen of water.

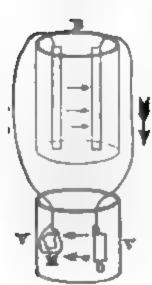
487. It will be perceived, that there is a great analogy between the acm of a simple battery for producing metallic depositions, and the constant strery of Professor Daniell. It is to the substitution of copper for hydrom, as above mentioned, that the superiority of Daniell's battery is to be cribed. By this change, cohesion between the cathion (360) and the suron of the conductor, acting as the copper plate in a galvanic triad (\$13, c.), is prevented; and this conductor, incressntly renovated superficially,

ments always a bright metallic surface to the liquid.

488. After a constant battery, of the construction thus described, has en in operation for some time, on removing the coating of metallic copper am the surface of the copper plate, it is found to constitute a sheet of the posited metal of greater or less thickness, and which is a perfect fac mile of the surface on which it has been formed, excepting that depresme and elevations are represented in the new plate, in the opposite kind Although these results could not have escaped the observation of erators with the constant battery, no useful consequences flowed from the at which they were of a nature to afford, until the year 1888, when, rough the ingenuity of Mr. Spencer, of Liverpool, and of Mr. Jacoby, of Petersburg, the new and important arts of electrotype and electro-gilding tre invented.

489. If a piece of coin\* be made the negative surface of a simple gal-





\* Figure 25 exemplifies a constant battery, arranged so as to serve for the electrotype process. A plate of sinc, Z, is suspended in diluted sulphurio acid, on one side of the porous duphragm, a medal on the other, being severally

attached to the ends of a copper wire.

Figure 96 will convey an idea of another mode in some respects preferable. The mould M is connected by means of a wire with the zinc of a Daniell's constant battery, a sail of copper is connected by the same means with the copper plate. Thus arranged, the coil and mould, or medal, acting as electrodes, are plunged into a concentrated seletion of sulphate of copper acidulated by sulphuric acid, con-tained in a vessel without any porous partition, the sail being made to enter the solution first. It has been montioned, that the true sinc end of a galvanic battery is nega tive. Of course the metal communicating with the zine becomes the negative pole or cathode of a galvanic tried, (358) while the copper coil becomes the positive electrode. Under these circumstances the copper coil is exidized and dissolved by the acid, while an equivalent portion of this metal is deposited on the mould.

The roll of sheet copper should be introduced first into

the solution and must not touch the mould. Great precautions as to cleanliness must be employed in this, as well as in the former method of manipulating. When an arrangement of this description is set in action, for each equivalent of sinc, dissolved in the battery, an equivalent of sepper is dissolved from the coil of sheet copper, and is prevanic battery, by attaching it to a piece of zinc by means of a short az d copper wire, the zinc being acted upon by a saline or acid solution on one side of a porous diaphragm, while the coin is plunged into a mice a sulphate of copper upon the other side, the coin will be speed a ... = with a light rose-coloured film of reduced copper, which by contact to action for some hours, will become proportionably thicker, and may in be separated from the coin, by the application of a moderate heat. Is metallic plate thus obtained, as already mentioned, will display a proreverse impression of the com, and by making the copy thus promine a conducting surface of a similar arrangement, a perfect impressor at lief of the original will result. The conducting wire, and all that pe tions of the coin upon which copper is not to be deposited, must be one or varnished, with a non-conducting substance. The surface to be case must be perfectly free from uncleanliness, from grease especially. The := should be immersed in the acid a little before the coin is plunged at 2 metallic solution, and the surface of the last should then be freed in a wbubbles by means of a feather or glass rod. The rapidity with which is deposition is effected, is greater in proportion, as the temperature of its solution is higher. But there is a certain degree of speed which would not be exceeded, as the texture of the metallic deposition is more compaswhen the process is slow.

490. A more expeditious method of producing the copy of a media a relief, consists in first making a copy, by stamping clean sheet lead upon 5 or pressing it upon fusible metal, while sufficiently soft: or taking at a pression on plaster, the surface of the cast thus obtained, being made > conduct by means of a film of plumbago. The impressions thus process

may be substituted for the medal in the process above described.

491. These last mentioned expedients are peculiarly appropriate a

cipitated upon the surface of the mould connected with the gine plate. posing between the zinc and copper of the general ng cell a number of decorate cells, similar to that already described the moula of one being a number of second copper roll of the next, and so on in the same order, any number of meanly of may be copied at once and at the expense of but a single equivalent to a

The wires connected with the rolls of sheet copper should be a arm she the their being dissolved off the moulds, and the copper electrodes a noble or him. tically and parallel to each other in the solution, and this last should be seen stirred with a glass rod to preserve an uniform degree of saturation tile agree mass of liquid

Any conducting surface, negative to the zine of the lattery cell may be well upon in this apparatus. Hence, if surfaces of wax, wood, paster & receive a thin coating of plu magn, by rabbing that material in the brush, they may be made to receive a Coating of metallic copper very mar r to

form and as perfect as if they had been then selves of metal

The accuracy with which the minutest rievalt ins and depressions of a surface may be copied by means of the electrotype process, as so great the method has even been applied to the multiplication of Daguerreotype pictures at the copies thus produced, are in every respect equal to the originals in perfect finish and minuteness of execution. They possess, besides, the advantage of the indestructible by friction and slight mechanical injuries. For this beaut tallage to tion of the discovery of Spencer and Jacobi, we are indebted to Mr Eng. :

The above described processes have not been confined to the production of the trotypes. They have been recently applied by M. Becquerel, in France and "Jacobi, in Russia, to the separation of the precious metals from their ores, and agree ably to a statement made in some of the foreign papers, a mass of gold deposited a this manner, weighing half a pound, was presented a few months since to the K.

of Russia, by M. Jacobi

copying engravings on copper, or wood, or in procuring stereotype plates from printing type set up so as to be ready for the press.

#### Of Electro-gilding and Electro-plating, &c.

492. But a more generally useful application of the principle is the electro-gilding and plating, now in almost universal use. If any polished erticle of steel, copper, or brass, be connected with the zinc of the bettery, and placed opposite a fine wire of gold or platinum, connected with the copper of the same, and if the two electrodes thus formed be plunged into a very dilute solution of chloride of gold, or chloride of silver, to which an equivalent of cyanide of potassium in solution has been added, a deposit of gold or silver will be effected, and the article will receive a fine coherent film of gold or silver, with a polish equal to that of the surface on which the deposit is made. Platinum may also be worked in the same way, though the operation is very far from being uniformly successful, and further experiments are wanting to render the process less precarious.

493. The process of electro-gilding has been employed in England, by Prof. Wheatston, and in Germany, by Prof. Steinheil, for the protection of the reflecting surfaces of the specula of reflecting telescopes, and specula of good instruments may be multiplied indefinitely, as well as very cheaply, by taking electrotype casts of their surfaces, and then gilding the reflecting

surfaces of the copper specula thus obtained.

494. In depositing gold, silver and platina, by the electrotype process, it is necessary to employ currents of feeble intensity, and very weak metallic solutions, as the metals are very liable under the action of a very energetic current to be precipitated in a pulverulent form. By Mr. Davis, of Boston, the current of the magneto-electric machine was first employed for this purpose, and a patent has recently been obtained in England for the same process and manipulations.\*

495. Besides the applications of the electrotype which we have noticed, many others have been contemplated or employed, of which a due regard

to brevity renders it inexpedient to give the details in this treatise.

#### Of Metallo-Chromes.†

496. The name which forms the preceding head, has been given to certain beautiful colours, which may be produced upon a plate of highly polinhed steel by the electrolyzation of acctate of lead, while in contact with the steel surface. The following directions are given for the production of the phenomenon in question. Place the steel plate in a glass basin contain-

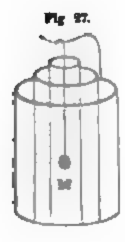


Fig 27 represents an apparatus suitable either for the process. of electro-silvering, or that of electro-gilding. It consists of a jar of glass, or glased porcelain, within which is situated a jar of porous porcelain containing the metallic solution (492). The space between the two jars is occupied by a cylinder of sinc, and sulphutic acid very much diluted. A wire proceeds from the zinc cylinder of sinc, and sulphutic acid very much diluted. der, as represented in the engraving, for the purpose of supporting the article which is to receive the metallic coating. It is recommended to dip the article, which is the object of the process, in nitric acid, just before placing it in the circuit.

† The etymology of this name from  $\chi_{equi}$ , colour, in Greek, must be sufficiently obvious. As designating an iridescence preduced by the peroxide of lead, it seems to me that a more apprepriate appellation would be that of plumboxychromes.

which some regular device has been cut. A small rim of wood should placed over the card, and upon that a copper disk. On contact brog and from five to twenty degrees, in a circuit formed with two or three card, and anall constant battery, the steel plate being made negative, the opper in positive, the deposit will be effected, and a series of exquisite colors of adorn the steel plate. These variegated tints are produced by the man tenuities of films of highly oxidized lead. Two sets of presente colors are produced, complimentary to each other, one by transmitted land the other by that which is reflected from the steel. The colours are depicted in the greatest perfection, by placing the plate before a window, and or clining over it a sheet of white paper, at an angle of forty-five degrees.

# Of the Galvanic Powers and Effects of Animal Organ

497. Under the preceding designation, it seems proper to treat of the galvanic apparatus and phenomena, with owe their existence to organic masses. It will be preceived by the student, who has given due attention to the preceding pages of this treatise, that this topic, has necessarily been in some degree anticipated. The reference made to the convulsions produced in frogs, by an inherent electrical power, and to the shocks produced by electrical fishes, has already introduced organic galvanism to be notice.

498. Evidently of all the sources of the galvano-electric current, none are more interesting than the organs with which nature has furnished the torpedo, the gymnotus and other electrical fishes.

499. Of the Torpedo.—Experimental evidence of the identity of the cause of the miraculous power of the torpedo, in giving shocks to other animals, seems first to have been satisfactorily obtained in the year 1792, through the exertions of Walsh, a sagacrous electrician. In the following abstracts from a letter addressed to Dr. Franklathis able observer gives an account of the facts which he had ascertained.

500. "It is with peculiar satisfaction, that I make to you my first communication, that the effect of the torpedo appears to be absolutely electron. We have observed that the back and breast of the animal are in different electrical states, \* \* \* we have been enabled to direct his shocks through a circuit of four persons, all feeling them; likewise through a considerance extent of wire, held by two insulated persons, one touching the back, and the other the breast of the fish. These experiments have been varied a many ways, and repeated times without number, and they all determines the choice of conductors to be the same in the torpedo as in the Leyder phial."

501. Respecting the torpedo, I will quote a few paragraphs from a report made to the Academy of Sciences at Paris.

502. "The sensation which the torpedo causes when it is touched, has long ago attracted the attention of physicists and physiologists, on account of its analogy with that produced by an electrical battery, but it is only a few years since that it has been decidedly proved that both were owing to the same cause. Although all the principal circumstances of this phenomenon had previously been carefully studied, yet no one had succeeded in de-

monstrating its electrical origin from the want of suitable apparatus.

503. "John Davy made known, in a paper published in 1832, a great number of important data, such as the action of the discharge upon the magnetic needle, and chemical compounds; but the direction of the electrical current produced on this occasion was not well known until after the experiments made at Venice, 1835, by two of your members, and from which it resulted that the superior part of the electrical organ gives positive electricity, and the inferior part negative electricity. Matteucci has confirmed, with the galvanometer and frogs prepared after the method of Galvani, the observations which we had made respecting this point, as well as others also relating to the torpedo, for which we are indebted to various philosophers; at the same time he has demonstrated some new facts, of which the following is a short account.

504. "He commences by showing that when the torpedo lances its discharge, no change of volume is observed in its body. When the animal is . possessed of great liveliness the sensation is felt at whatever point of the body it may be touched, but when its vitality is considerably diminished, the discharge is no longer felt, except by touching the electrical organs at two

different points.

- 505. "Matteucci establishes the general laws of the distribution of electricity in this manner:—1. All the points of the dorsal part of the organ are positive relatively to the points of the ventral part; a fact already known. 2. The points of the organ on the dorsal surface placed above the nerves which enter it are positive in respect to the other points of the same dorsal surface. 8. The points of the organ situated on the ventral surface corresponding to the points which are positive on the dorsal surface, are negative in respect to the other points of the ventral surface. 4. The intensity of the current varies with the extent of the platina plates which terminate the galvanometer, and with which the two surfaces of the organ are touched.
- 506. "When the torpedo is very excitable the current may be compared to that of a pile consisting of a great number of pairs charged with a good conducting active liquid; whilst, on the other hand, when its liveliness is weak, the electric current resembles that of a pile composed of a small number of elements.
- 507. "The spark which accompanies the discharge in the electrical fishes was remarked for the first time by Walsch in the Gymnotis; many vain efforts have been made since to reproduce it; MM. Matteucci and Limari have succeeded in obtaining it in every case from the torpedo; both these philosophers claim the priority of the observation.

508. "Matteucci has since succeeded in obtaining the spark by placing

the torpedo upon an isolated plate of metal, and placing another photometal above it, then fixing to each of them a gold leaf separated the corine the other by the distance of half a millimetre. By slightly normal upper metallic plate the animal became irritated, and at the same count the two leaves approached one another and the report of the spart was

stantly heard.

of the electrical organs, without detaching the epidermis, one of the polyment of the galvanometer was inserted in the organ near the outward of the plate was put in communication with one of the four near the needle deviated four degrees in the common direction of the doctaged the torpedo; on tying the nerves there was no longer any deviation.

result appears to us very remarkable.

from the want of torpedoes, go to prove, 1st, that the electricity which per duces the discharge proceeds from the last lobe of the brain, and a mitted by the nerves to the organ; 2, that the discharge coasing vote influence of the electric current, when the nerves are tied, must, in ordat be transmitted, find in the nerve a particular molecular disposition, and clusion to which the electro-physiological phenomena of the frog real lead, as one of us (M. Becquerel) has indicated in various places in the

treatise on electricity.

the contact of two different metals in communication with the most of nerves of a frog sufficed to make it contract, the experiments have writed infinitely in the hope to discover in this phenomenon the cause with constitutes life in animated bodies. The most remarkable fact, for the we are also indebted to Galvani, is that which relates to the contract produced by the simple contact of the muscles and nerves without the mediary of metallic armatures. It is now nearly demonstrated that action does not proceed from a chemical action, but from the interest of rent of the frog, which has been indicated with so much sagain to Nobili."

- 512. The opinion and facts with which these quotates tend to make the reader acquainted, seem to me to prose that if Volta was right in conceiving the powers of his per as independent of vital action, still Galvani was justified a supposing that animal organization could give rise to the logous results.
- 513. Of the Gymnotus Electricus.—According to Hopboldt, the gymnotus abounds in the small rivers which is into the Orinoco. It employs its electrical powers as other animals do their horns, teeth, or fangs, to secure its into or repel aggression. Hence, horses or mules are attacked by them, when forced reluctantly to enter the streams of which they reside; and while, on the one hand, the quadrupeds suffer severely, from the shocks which they recent

their assailants are so exhausted, as to be unable either to

give further shocks, or to escape from seizure.

514. The gymnotus is sometimes five feet in length. One of the length of forty inches was lately conveyed to England, and put at the disposal of Faraday, who has given a most valuable account of the various phenomena noticed by himself, and other intelligent observers.

515. The shock was most powerful when the hands were made, the one to touch the animal near the head, the other near the tail, and was less severe in proportion as the distance between the hands was less. When the points of contact were in a line at right angles to the spine, little sensation could be produced.

516. The farther the hands were from the fish, while the ine extending from one to the other was parallel to the spine, the less was the sensation experienced, and the diminution became greater in proportion, as the line joining he points of immersion deviated from parallelism with the mpine.

517. Conductors of an ingenious construction, being ap-slied to the anterior and posterior parts of the gymnotus, Faraday gives an account of the results in the following

words:--

518. "A galvanometer was readily affected. It was not particularly belicate; for zinc and platina plates on the upper and lower surface of the ongue did not cause a permanent deflection of more than 25°; yet when he fish gave a powerful discharge the deflection was as much as 80°, and n one case even 40°. The deflection was constantly in a given direction, he electric current being always from the anterior parts of the animal brough the galvanometer wire to the posterior parts. The former were herefore for the time externally positive, and the latter negative.

519. "Making a magnet. When a little helix containing twenty-two bet of silked wire wound on a quill was put into the circuit, and an ansealed steel needle placed in the helix, the needle became a magnet, and be direction of its polarity in every case indicated a current from the anprior to the posterior parts of the gymnotus through the conductors used.

520. "Chemical decomposition. Polar decomposition of a solution of odide of potassium was easily obtained. Three or four folds of paper noistened in the solution were placed between a platina plate and the end fa wire also of platina, these being respectively connected with the two addle conductors. Whenever the wire was in conjunction with the conuctor at the fore part of the gymnotus, iodine appeared at its extremity: at when connected with the other conductor none was evolved at the place n the paper where it before appeared. So that here again the direction of he current proved to be the same as that given by the former tests.

521. "By this test I compared the mide tions before and behind it, and found the applied to the middle was negative to the rior parts, was, on the contrary, positive near the tail. So that within certain limit ternally at the time of the shock appears negative to other parts anterior to it, and

522. "Evolution of heat. Using a longing to Mr. Gassiot, we thought we w when the deflection of the galvanometer tion of temperature. I was not observin of those who at first believed they saw the

523. "Spark. The electric spark wanto-electric coil, with a core of soft iron to the end of one of the saddle collectors, file; another file was made fast to the enson then rubbed the point of one of these whilst another person put the collectors excite it to action. By the friction of the very frequently; and the object was to through the wire and belix, and by break make the electricity sensible as a spark.

524. "The spark was obtained four tinent saw it. That it was not due to the shown by its not occurring when the Bdently of the animal. Since then I have revolving steel plate, cut file fashion on it of iron, copper and silver, with all of who

525. It has been shown there may increase voltaic power, or quantity, the other, that of increase the latter, the conducting power the case of an animal created course, to secure to these ele weapon with which they are en an enormous quantity of the retheir disposal.

526. The shock from the gyn observations were made, was, a to that of a large Leyden batter of intensity; or to a voltaic ser the circuit should be completed

527. He conceived it to cor

<sup>&</sup>quot; In more recent experiments of the same to At a later meeting, at which attempts gold leaves, the spark was obtained directly coil being removed, and only short wires (by



∄i

- :city equivalent to a high charge from a coated surface of :3500 square inches.

528. The gymnotus gives sometimes two, and at other times, even three successive shocks; as had been previously noticed in the case of the torpedo, by various distinguished observers.

529. At the moment when the fish wills the shock, of two parts of the body parallel to the spine, that which is

nearer the head will be positive, the other negative.

530. The fish has four electric organs, which, it is conceived by Faraday, it may have the power to throw into action separately, or together, and so as, to a certain extent, to direct the shock at the moment of the discharge. But it is not inferred that the direction of the current can be controlled after it has entered the liquid or other surrounding bodies.

531. I saw the gymnotus with which these experiments were made in 1841, and received a shock from it. I was moreover gratified at seeing what I had not been led to expect, the deflagration of a thin metallic leaf by subjec-

tion to a circuit formed with the aid of the animal.

- 532. The gymnotus can stun or kill fish in various positions of its body, but in killing a small fish, thrown into the tub, was found to coil i self so as to make the fish occupy the situation of a diameter to the resulting coil. In an instant the fish was struck dead and motionless, and turning on its side, was soon bolted by its hungry assailant.
- 533. "Living (says Faraday) in the midst of such a good conductor as water, the first thoughts are thoughts of surprise that it can sensibly electrify any thing; but a little consideration soon makes one conscious of many points of great beauty, illustrating the wisdom of the whole arrangement. Thus the very conducting power which the water has; that which it gives to the moistened skin of the fish or animal to be struck; the extent of surface by which the fish and the water conducting the charge to it are in contact; all conduce to favour and increase the shock upon the doomed animal, and are in the most perfect contrast with the inefficient state of things which would exist if the gymnotus and the fish were surrounded by air; and at the same time that the power is one of low intensity, so that a dry skin wards it off, though a moist one conducts it; so is it one of great quantity, that though the surrounding water does conduct away much, enough to produce a full effect may take its course through the body of the fish that is to be caught for food, or the enemy that is to be conquered.

534. "Another remarkable result of the relation of the gymnotus and its prey to the medium around them is, that the larger the fish to be killed or

stunned, the greater will be the shock to which it is subject, thout to gymnotus may exert only an equal power; for the large fish has much through its body those currents of electricity, which, in the case of a call one, would have been conveyed harmless by the water at its sides.

animal, being made conscious of it, probably, by the mechanical units he receives, caused by the spasms into which it is thrown. When I to the him with my hands, he gave me shock after shock; but when I to the him with glass rods, or the insulated conductors, he gave one or two shot, felt by others having their hands in at a distance, but then crased to me the influence, as if made aware it had not the desired effect. Again, the has been touched with the conductors several times, for expension the galvanometer or other apparatus, and appears to be languad or mainent, and not willing to give shocks, yet being touched by the hands, they by convulsive motion, have informed him that a sensitive thing was provided and he has quickly shown his power and his willingness to astomath experimenter.

organs of the torpedo, gymnotus, and similar fishes, cannot be constant as essentially connected with those which are of high and direct importance to the life of the animal, but to belong rather to the common tendents; and it has also been found that such torpedoes as have been deposed of the use of their peculiar organs, have continued the functions of he can as well as those in which they were allowed to remain. These, with the considerations, lead me to look at these parts with a hope that they upon close investigation prove to be a species of natural apparatus, by the of which we may apply the principles of action and reaction in the investigation in the investigation prove to be a species of natural apparatus, by the of which we may apply the principles of action and reaction in the investigation.

gation of the nature of the nervous influence."

537. I cannot perceive, with Faraday, that the facts above mentioned, as stated by him, are all consistent with the idea of the powers of the gymnotus, being due to a great quantity of statical electricity of low intensity.

538. The fact which appears to me inexplicable, upon this view of the subject is, that a shock should be received by a person having only one hand in the water containing

the fish.

539. Agreeably to universal experience, no shock can be received by any person who is not within the electrical circuit through which the discharge is made; or of some other induced circuit in the vicinity. But within what circuit can a single hand be brought, unless the body and legs of the person to whom it belongs, the floor on which he stands, and the tub containing the gymnotus, be made to form a conducting communication between the organic electrodes of that animal. But if the electricity discharged through these electrodes be of an intensity too low for

nem to discharge themselves through the water, a fortiori, must be too low to make its way by means of the floor f which the conducting power must be vastly inferior to hat of water.

540. I regret that it was not ascertained whether this esult could not have been prevented by placing the tub

pon glass legs, as dry as possible.

541. I doubt if a voltaic series of 100 large pairs, placed n water as the fish was, would cause shocks to a person

on immersing only one hand.

542. A battery of 100 pairs, when out of water, will not, agreeably to my experience, give a shock to one hand. Of course, it could not do more when submerged in that liquid.

543. It is difficult to reconcile this fact with the idea, that the powers of the gymnotus are those of a large Leyden battery lowly charged, unless it be supposed that the animal is enabled to project the electric fluid in obedience to its will, as the porcupine has been supposed to

shoot forth his quills.

544. The anatomical researches of Hunter, and other anatomists, have been considered as demonstrating, that the electrical organs of the torpedo, bear a very striking resemblance to a galvanic series in the columnar forms of the original pile of Volta. They have been alleged to consist of hexagonal columns, amounting in one instance, to one thousand one hundred and sixty-two, arranged like the cells of a honey comb, each column consisting of a pile of flat disks separated from each other by cellular tissue. The organ is abundantly furnished with nerves. Nevertheless, little satisfaction seems to have been afforded by the investigation of anatomists respecting the mechanism, by which nature has endowed this animal and other electrical fishes with their peculiar powers.

545. Both Matteucci and Schoenbien concur in considering, that neither dissections, observation, nor experimental inquiry, have thus far afforded any competent explanation of the apparatus by which the powers of the torpedo or gymnotus are imparted and made obedient to animal will. By the former it is alleged, "that it is impossible to find the least analogy between piles, secondary coils, batteries, and the electric organ of the torpedo:"

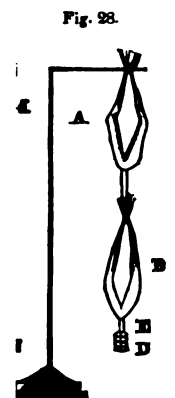
of Faraday, with the same gymnotus, it is stated as result of his mature consideration, "that the true can the phenomena is still completely obscure, and must and be sought for in the physical or chemical constitution, in a fixed organization of certain parts of the animal; that there exists, without our being able at present to termine how, an intimate connexion between the vactions dependent on the will of the fish, and the physical or chemical constitution.

#### Of the Observations of Nobili, Matteucci, and Others.

546. It has been mentioned that Galvani considered that the courd produced in frogs by contact with metals, or the proximity of the duch from the conductor of an electrical machine, as more or less depends a diversity of electrical excitement in the different animal organs.

547. That Volta having subsequently discovered his pile to be a of producing effects altogether independent of the animal, similar is character and vastly more powerful than those noticed by Galran eminent philosopher was led into an error, which was generally adea supposing that the organs of a frog, when convulsed by an closur charge, are in all cases the medium, not the source of the current. I recent observations of Nobili and Matteucci, have shown that although impressions of Galvani may not have been warranted by any of his experiments, they are fully justified by those which have been to made.

548. From the experiments of Matteucci, it appears, that in forge is a current, peculiar to them, proceeding from the feet towards the This can be made sensible either by the deviation caused in the need very delicate Sweiger's galvanometer of 2500 circumvolutions, or b contractions produced in the leg of another frog, of which the da spinal nerve is made the medium of discharge. I shall call a fing's and nerve thus prepared, a galvanoscopic frog limb. To produce the rent in question, only the leg of the animal is necessary; althou several experiments, the leg, thigh, denuded nerve, and a portion of spine, are mentioned as being employed. The author in one of his maries, uses the following language: "Each member of a frog, prising the leg, thigh, spinal nerve, and a small portion of the spine stitutes a complete electromotor." Yet it fully appears from his subsenarrative, that the thigh, nerve, and spine, contribute nothing to the tromotive energy. The nerve, from its low conducting power, it diminishes than increases the intensity of the discharge. It follows the statement, above made, should have been reserved for the leg of frog; which, agreeably to the author's observations, is really an elect. tor as much as any galvanic triad (312). It does not appear to be mined, what portions of the leg perform respectively the parts of c and zinc: the sangumeous liquid, retained in the pores of the fiesh, set act as the electrolyte-



549. Nobili had found, that by disposing a number of frogs in such order as to resemble a voltaic series, an increase of power in the resulting electrical current would ensue. To construct such a series, it is sufficient that the frogs, duly prepared, be supported by one or more non-conductors, so that their dissimilar extremities may be in contact, or communicate through weak brine, forming thus a species of couronne des The adjoining figure illustrates one convenient mode among others, by which the frog limbs may be connected, so as to be in voltaic order. Of course the series might consist of several similarly associated. M N is a glass rod, by which the frog's limbs, A, and those at B, are supported and insulated. According to Sturgeon, when the spine, D, of B, is, by means of another glass rod, brought into contact with the feet of A, convulsions are produced. Moreover, it will be

en, that agreeably to the experiments of Nobili, and those of Matteucci, nen a conducting communication is made between the extremities of such organic series, through the coil of a galvanometer of 2500 circumvoluas, deviations are produced, which are, as far as tried, greater in propor-

**n** to the number of the series.

550. At first the frogs thus employed were prepared as above described, as to consist of the legs, thighs, and a bit of spine, but Matteucci discored that, as naturally associated, one limb operated in some degree as a charger to the other, so that on the removal of one, the other became ore efficient. It has been mentioned that the thigh and nerve added noing to the electromotive power. Accordingly it was found that a battery ight be made of legs only, the feet belonging to one leg touching the nuded muscular extremity of another, so that the dissimilar parts of the bole number were in contact in voltaic order (362).

551. Of the voltaic series thus formed, the effects upon the needle of the lvanometer increased within certain limits with the number, so that when e element could produce a deviation of only four or five degrees, the ses would produce fifteen to twenty degrees.\* These deviations were of a iture to show that the current sets invariably from the feet towards the st of the limb. The current, thus described, is designated by Matteucci the current proper to the frog. He has not been enabled to detect any ing analogous to this current in any other animals of a neighbouring

ade, such as lizards, tortoises, or cels.

552. Nevertheless, agreeably to the same authority, currents may be own to take place from the internal to the external portions of the muses of both warm and cold blooded animals, by which, deviations in the dvanometrical needle and contractions in a galvanoscopic frog's leg may produced, as striking as those resulting from the current proper to ogs.

553. Sections of thighs of frogs, of the bodies of eels, or of the thighs of bbits or pigeons, being arranged in a series of eight or ten on a varnished

ank, so that the internal surface of one should touch the external surface

<sup>\*</sup> The deviations of the needle of a galvanometer are estimated by a semi-circle, advated to ninety degrees on each side of the point at which it rests when undisrbed.



the external, the skin being removed, a current is c parts to the outer.

556. That this current varies as to intensity

whose muscles are employed.

557: That by disposing duly made sections of described, in a series, voltaically, the intensity of t tion to the number of the sections so associated.

558. The author adverts to a fact which he had when water and blood communicate through a liqu is induced from the latter to the former. Foreses presented as accounting for the current from the in mal muscle, he points out that a current, resulting the muscle and the water used as a conductor, its effect on the needle to that observed, which is of the direction being from the water towards the sale conclusive answer to the objection thus founded, the galvanometer were due to the electromotive water, arranging the sections, so as to form a voltal augmented the intensity as it was actually ascertain

559. Further, it is alleged that the current detebetween water and blood, endures without diminithat which takes place in the muscles of recently sient, and ceases sooner in proportion as the anima

560. Contractions were produced in the leg of when the nerve was made to complete the poles of of moistened paper. Even when the paper was so that the galvanometrical needle was not affected by platina blades in contact with the paper at different communications between the paper and frog, cor longer the strip of paper employed for a part of the number of the sections requisite to produce the con-

Of Galvano-Electrical Currents in Lix

An experiment is described, in which several live frogs were subrected to a process, which may be literally considered as excruciating, by
iling their feet against a varnished plank, removing their legs, skinning
reir thighs, and in cutting in half each alternate thigh; then the uncut
righ of the first, and the raw surface of the remaining half of the next
righ, were brought into contact in voltaic succession. By these, a series
four elements gave a current of twelve degrees, always from the interior
the surface.

563. The galvanometrical deviations caused by a current obtained from z live sheep, or rabbit, by the means above described (560), was more san six times greater than those obtained from a frog in like manner.

564. The potency of muscular batteries, in which the size of each element in one, was uniformly double or triple that of those in the other, was sund nearly the same; since, when they were so arranged as that the one ounteracted the other, there was scarcely any current produced by the xcess of power in the larger element. When, of two batteries of frog aighs, of the same size and number, the nerves were removed only from me, no diversity of power was found to ensue: so that the muscular curent depends neither as to quantity, nor as to intensity, on the integrity of he nervous system of motion and sensorial power. Two batteries of ten stements each, being constructed, the one of the thighs of frogs, of which he spinal marrow had been destroyed by a hot iron, and their lower limbs thus paralyzed, the other differing only in being made from animals of the mme kind, in which the spine had sustained no injury, it was found that where the two batteries were made to act in opposition in the same circuit completed by the galvanometor, the deviations of the needle in the direction of the former, was from sixteen to eighteen degrees. In fact, this battery gave separately, from fifty to fifty-five, while the other gave only from forty to forty-five degrees.

565. This is considered as confirming the inference already made, that the integrity of the motive and sensorial nervous system, has no bearing on the ntensity or the direction of galvano-muscular currents. Whether or not, the frogs from which batteries were constructed, had been rendered drowsy by pium, nux vomica, or other narcotic poisons, made no difference in the surrents which they gave; excepting in one instance, in which it seemed is if a feeble dose augmented the power of the current. Moreover, it is not to be inferred, that the capacity to produce a current, is not enfeebled or destroyed, when narcotics have been used to an extent to produce death. Frogs, thus poisoned, are as unsuitable for a battery, as those which have been dead for a much longer time, after being killed and prepared in the asual manner. In operating with the muscles of a pigeon, which had been more or less poisoned, similar results were attained. Contrasting the galvano-electric power of equal series of the muscles of frogs or pigeons killed by immersion in carbonic acid, cyanhydric acid, or arsenuretted hydrogen, and of such as were killed as usual, no superiority was found in any case, excepting that a little superiority appeared in the muscles of those which had been poisoned. Yet, in the case of muscles from frogs, or pigeons, killed by sulphuretted hydrogen, a great loss of power was discovered, as contrasted with the muscles of animals prepared in the ordinary way; since, when by a similar series of the latter, scarcely any deviation could be obtained, by the former fifteen degrees were effected.

The following is the author's summary of his inferences.

566. 1st. The intensity of the galvano-muscular current, as measured

by deviations of the galvanometrical needle, varies in cold bloods are proportionally to the temperature of the medium in which they have for some time previously to the experiments.

587. The durability of the current, is less in proportion, as the man

more elevated in the scale of beings.

568. The intensity of the current varies as the animal has been too worse fed, and is invigorated by inflammation, or sangumous count resulting from a wound.

569. It is independent of the integrity of the nervous system, as many

the sensorial and motive power.

570. Narcotic poisons have scarcely any influence on this current

spects intensity, and as to its direction they have none-

571. Among gaseous poisons, sulphuretted hydrogen, is to a to extent, productive of an enfeebling influence, while cyanhydro acd, a seniuretted hydrogen, if productive of any change, strengthen a little.

572. The direction and the strength of the galvano-muscular culture altogether independent of any assistance from the nerves. who only as imperfect conductors, representing and transmitting the electrical conductors.

citement of the muscle with which they are associated.

573. Messrs. Pacinotti and Puccinotti, of Pisa, in a great may a ments, in which one of the blades of a galvanometer was plunged at leg, the other into the brain, obtained deviations of the needle, which at the same time sufficiently striking, and of a nature to demonstrate existence of a current flowing from the brain towards the muscular pathe animal fabric.

avoiding erroneous results. He suggests that the action of the blood avoiding erroneous results. He suggests that the action of the blood flows copiously from the wound in the brain as usually made, is all by its reaction with the terminating blades of the galvanometer, to predeviations, especially as, in order not to diminish the vitality of the method been preferred to perforate the head the last. Merely the first sulting from moving the blades is sufficient to increase the deviations having himself taken various precautions, the deviation on first new was always such as to comport with a current in the direction in brain towards the feet, as observed by the electricians above named. Matteucci's statement, that deviation varied from eight to ten degree as the current did not persist in its direction after the first pour very much enfectled after the third immersion, and was very every much enfectled after the third immersion, and was very eversed, it does not seem that much rehance is to be placed on the case and direction of the current in question.

575. Assuming, however, that there is a current from the brant is the lower parts of the body or limbs, our author conceives, that it is a sequence of the nervous system acting as a conductor to the mis masses into which the nerves rannity. Yet, he admits, that this is a altogether compatible with the fact of the production of a current cither by a contact with the interior of any of the muscles, or with a the muscular surface; and, further investigation is requisite for the c'.

tion of this subject.

576. The author concludes with this observation. Perhaps the a system may exercise on the galvano-animal current an indirect not arising from its participation in the process of muscular nutrition, we is not, however, possible to prove by direct experiments.

EXHIBITION OF VOLTAIC SERIES OF VARIOUS CONSTRUCTIONS.

- 577. Voltaic Pile, as originally constructed, exemplified—also the Couronne des Tasses.
- 578. Apparatus, employed by Sir Humphrey Davy, exemplified by two porcelain troughs, in each of which are suspended ten pairs, consisting severally of a zinc and a copper plate.
- 579. Apparatus exhibited, in which the cells are made by partitions of glass, and the galvanic pairs, instead of being attached to beams in order to be lifted out of the acid, are kept permanently in their cells; the acid being removed by a partial revolution of the trough upon pivots, when the action is to be suspended.
- 580. Construction of Cruikshank pairs and trough illustrated.
- 581. Galvanic deflagrators exhibited of various dimensions and numerical extension, combining the advantages of the Cruikshank trough with those of the deflagrator.
  - 582. Effect of the circuit upon the animal frame.
  - 583. Iodine evolved and made evident by starch.
  - 584. Water decomposed, and recomposed.
- 585. Various apparatus for different amalgams proluced.
- 586. Deflagration, fusion, and volatilization of charcoal, netals, anthracite, and plumbago, exhibited. Likewise of rarious metallic wires, of tin-foil, tinsel, and mercury. Fusion and incorporation of iron and platina, under water. Hydrate of potash deflagrated upon charcoal, and upon a piece of silver coin.
- 587. Fusion of platina, on contact with chloride of calcium and other solutions in the voltaic circuit.
- 588. De Luc's columnar apparatus exhibited and explained.

# Exhibition of

Grove's gas battery.

Daniell's original constant battery.

Improved constant battery. Grove's sustaining battery.

Apparatus and processes for electro-typing, also in electro-gilding and silvering, &c., and metallo chrome.

# Of a Process for Rock-blasting.

589. I have already adverted to the efficacious perfectly safe process, for rock-blasting and blowing " wrecks of vessels, fortifications, &c., with which I on trived, in 1831, to ignite twelve charges of guapowders the distance of one hundred and fifty feet (447). Hope that it will serve the cause of humanity to make this pill cess better known, I subjoin the original account of #1 published in the Journal of the Franklin Institute in 153 Vol. XX. page 221.

Description of a Process and an Apparatus for Blasting Rocks, by means of Gon Ignition. Communicated by Robert Hare, M.D., Professor of Chemistry in the vernty of Pennsylvania

590 I have observed various accounts in the newspapers of workmen tall

dreadfully lacerated, by the blasting of rocks

591. I have ascertained that by a new application of galvanteen, rocks my riven with less danger than that which attends the firing of a postal. I was min to attempt this improvement in consequence of an application by a patents Moses Shaw,) for assistance in perfecting his patented mode of blacking meta, by

electrical discharge from a Loydon jar.
532. In a letter dated June 1st, 1c31, he says, "I have been engaged to him rocks by means of a fulminating powder, introduced into several cavities, and use in all of them simultaneously, by a spark from an electrical manner means masses of a much larger size, and of a much more suitable size. object in view, may be produced, than by the old plan. I have, however to my inability to succeed in this method of blasting, during a great part of when, in consequence of the unfavourable state of the weather, the ign to a be effected by electricity in any mode which I have devised, or which has gested by others, although I have consulted all the best informed profess is to a large had access "

593. It occurred to me, as soon as this statement was made by Mr. Shaw that ignition of gunpowder, for the purposes he had in view, might be effected to vanie discharge from a deflagrator, or cal remotor in a mode which I have but in my eudiometrical experiments to ignite explosive giseous mixtures. This

cess is free from the uncertainty, which is always more or less attendant up-employment of mechanical electricity, for similar purposes.

594. The expectation thus arising, has since been fully verified. I have z as many as twelve charges of gunpowder at the distance of one hundred as a feet, from the galvanic machine employed. This distance is much greater a necessary to the safety of the operator, as the cuffigrator may be she ded a zto be injured by the explosion, and by mears of levers and pulleys it may be to act at any distance which may be preferred. There is no limit to the back charges which may be thus ignited, excepting those assigned, by economy, the size of the apparatus employed.

595 These remarks have reference to the principal and highly important the

Mr. Shaw a project, which is to ignite at once a great number of charges on as . perforations so drilled in a rock, as to cooperate simultaneously in the same personal by these means it is conceived that the stone may be separated into large prism or tabular masses, instead of being reduced to irregular fragments of an inferam of the common process of blicting by one charge, which renders that process

erfectly sufe.

606. This part of the subject I shall introduce by premising, that almost all the acidente which have taken place in blasting rocks, have occurred in one of the aree following modes .-

let. The explosion has taken place prematurely, before the operator has had time

s zetire.

2d. A premature explosion has ensued from a spark produced by the collision rising from ramming into the perforation, containing the powder, the brickdust, or and, or other matter, employed to close it.

3d. The fire not reaching the charge after the expiration of a period unusually mg, and the operator returning to accertain the cause of the supposed failure, an ag-lesion ensures when he is so near as to suffer by it, as in the instance near Nurristwn, published some years ago.

The means of communicating ignition, to which I have recorted, are as fol-

197 Three iron wires, of which one is of the smallest size need for wire gause, the there of the size (No. 24,) used by bottlers, are firmly twisted together. This is est accomplished by attaching them to the centre of the mandril of a lathe, which is mde to revolve while the other ends of the wires are held by a vice, so a s to keep som in a proper state of tension. After being thus twisted, a small pertion is unwisted, so as to get at and divide the larger wires by means of a pair of nippers. In this way the smaller wire is rendered the sole mean of metallic connexion between the larger ones. These are tied in a new kerf, so made in a small piece of agwood as to secure them from working, which, if permitted, would cause the mailer wire to break apart. At one end, the twist formed of the wires is soldered to be bettern of a ten take of a secure that a secure the mailer wire to break apart. maller were to break apart. At one end, the twist formed of the wires is soldered us to bottom of a un tube of a size to fill the perfuration in the rock to such a height a may be deemed proper. This tube being supplied with gunpowder, the crifice is lessed with a cork, perforated so that the twisted wire may pass out through it without touching the tube at any point above that where the finer portion alone interesses. To the outside of the tube, a copper wire, about No. 16, is soldered, long nough to extend to a stout copper wire proceeding from one of the poles of a galance deflagrator or calorimotor. The wire passing through the cork from the inside of the tube, is in like manner made to communicate with the other pole. The connexions between the wires and the rules, should be made by means of soft solder, are

the poles, should be made by means of soft solder, premously to which we must imagine that the tube has b introduced into a perforation made for its reception in a rock to be blasted. The tin tobe may be secured within the rock by the usual method of ramining in brickdust or sand, by means of a plug, having holes for the protection of the wires of communication already described."

506. The apparatus being thus prepared, by a galvanie discharge, produced by the movement of a lever through the quarter part of a circle, the finer wire is ignited, in the place where it intervenes solely in the circuit, so as to set

are to the surrounding gunpowder.

500. As the enclosure of the gunpowder in the tube, must render it impossible that it should be affected by a spark elicited by ramming, as no means of ignition can have access to the charge besides the galvanic discharge; and as this can only occur by design, without an intenti to commit murder or suicide, or by unpardonable neglect, it is inconceivable that an explosion can take place in this method of blasting, when any person is so situated as to

suffer by it.

600 It must be obvious that in all cases of blasting under water, the plan of a tin tube, and ignition by a galvanic

circuit, must be very eligible.

60). At A is represented a cylinder or tube of tinned iron, replete with gunpowder. At C, the twisted wires are represented as they protrude from the cylinder through a suck, by which the latter is closed at the upper end.

<sup>\*</sup> It has occurred to me that plaster of Paris might be used advantageously, as it reald require no reasoning, and might set with sufficient firmness.

The other ends of the wires are soldered to the metallic disk which form the solder of the cylinder. D represents the twisted wires as they appear when all the tap ones are cut, the smaller wire still uniting them. F represents the poor is greatedly prepared, and E the wires as when supported by the wood. The reason only to imagine the hole in the wood to be supplied with the foliminating control and covered by a fillet of paper or cloth, glued or pasted around the ways order to complete his conception of the wires as finally accounted and attention was

the cylinder A

602 Besides affording support to the larger wires, and thus protecting the main wire uniting them from fracture, the piece of dogwood which has been seemed means of the small hole represented in it, serves to hold, and to preserve to with the little wire, some fulminating powder. This not only farilities to expend ignition of the contents of the cylinder, but must make it expend mapped throughout the mass, and must, of course, cause it to be more ready to capital fractal metallic arsenic, and chlorate of potash finely powdered and mingled, make an cellent explosive powder for this purpose, being more ready to capital fractal and less so from other causes, than fulminating silver or more very. So pour sets used in lieu of arsenic. Yet the use of these is not necessary, as the purpose take fire directly from the wire, at least as effectually as in the usual mode. To mixture of gum phosphorus, sulphur and chlorate of potash used for markles, will answer, no doubt, in lieu of the preparation mentioned.



### **ELECTRO-MAGNETISM.**

720. The science of Electro-magnetism may be said to consist of a knowledge of a series of facts ascertained respecting the influence of active electric circuits with each other, or with bodies which may have been magnetized, or which are susceptible of being rendered magnetic. To describe and illustrate experimentally, some of the most striking among the facts, to which allusion is thus made; giving also such theoretic elucidation as the present state of science will afford, is all that the limits prescribed to this treatise will permit.

721. The science of Electro-magnetism may be considered as comprising the following branches, each producing, as its appropriate fruit, phenomena, which are not only highly wonderful and interesting, but likewise of great

practical importance.

#### SYNOPSIS.

Reciprocal reaction of Magnets.

Reaction between Magnets and Electrical Currents.

Attraction and Repulsion between Electrical Currents and the Movements thence arising.

Induction of Currents by Electrical Currents, or Electro-

dynamic Induction.

Induction of Magnetism by Electric Currents, or Electro-

magnetic Induction.

Induction of Electric Currents by Magnetism, or Magneto-

slectric Induction.

722. It had long been observed, that there were striking analogies, as well as discordancies, between the characteristics of electricity and magnetism. The similar poles of magnets, freely suspended, were observed to recede from each other, and dissimilar poles to approach each

<sup>\*</sup> Allusion is here made to what is called usually a magnetic needle, as represented in figure (35).

other, in a mode quite analogous to the separation proximation, under like circumstances, of substance

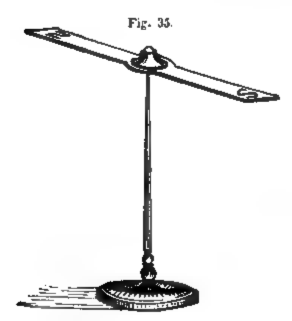
larly or dissimilarly electrified.

723. It had also been observed, that iron, vicinity of lightning rods, or otherwise indirectly a by lightning, had in some instances been rendered netic.

724. So far then, the existence of an analogy, or association, between electrical and magnetic ples was demonstrable; but, the magnetic repulsions a tractions took place under circumstances extreme favourable to the existence of the opposite elect The magnetic needle being a perfect conductor, the of course be no permanent existence of opposite of electrical excitement at the extremities. Moreover, pole of a magnet was more susceptible of being at by electrified bodies, than any other metallic wire, si suspended; nor did either of the electricities affe pole more than the other. The poles of the voltain ratus were found to be perfectly indifferent to the magnet, when it was presented to them, during tha ruption of the circuit, which was deemed necessary production of electrical excitement.

725. The contact of another magnetizable body of iron for instance, was found to strengthen and p the opposite magnetic excitement, while a similar a tion would be destructive of the opposite electric

ment.



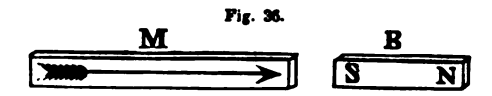
stances, then, it never of to any one, that the mipoles could be influent electricity, whether gaper or mechanical, till Properted, in the winter of ascertained the existent reciprocal influence, by the magnetic needle, wire connecting the polygalvanic apparatus. In ence to this influence, in

dently of terrestrial gravitation and magnetism, the when freely suspended, was found to assume invariant

more precisely, that of a tangent to a circle, concentric with, and at right angles to, the axis of the wire through which the current flows (742). Reference has been made to this wonderful reciprocal influence and the admirable galvanometrical invention to which it gave rise (306). It will be treated of more particularly under the appropriate head: Reaction between Magnets and Currents (731, &c.).

Of Magnetic Induction.

727. Magnetism has some analogy to statical electricity in the phenomena of induction, but utterly differs from it as respects conduction.



728. Thus the bar of iron B, figure 36, becomes magnetic, while in the vicinity of the permanent steel bar magnet M, precisely as B would be electrified by M, if previously charged by a machine: but, in the one case, there must be good insulation, while in the other none is requisite. The contact of M with B, would cause an electrical charge to be divided between them, whereas the same contact would cause the magnetic charge in B to be higher. In the one case the existing body loses power by contact, in the other it sustains no loss. That B is magnetized, may be shown by its attracting an iron nail or iron filings, if sufficiently near.

729. If several pieces of soft iron be placed successively in contact with each other, and with a steel magnet, as represented in the cut, figure 37, they will become magnets



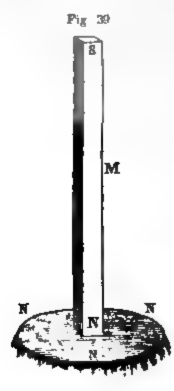
by induction, but will lose the property entirely if soon separated. Yet, a piece of iron kept for some time in a state of induced magnetism, retains the property to a certain extent, In proportion, as a steel bar is harder, it is less susceptible of receiving magnetism, but is more retentive of this property. On account of its superior retentive-

ness, all artificial magnets are made of steel, modernal hardened. Latterly our means of imparting magnetism is iron or steel have been immensely improved. A sai needle, or bar, may be permanently magnetized by drawing it, from one end to the other, over either poked a powerful magnet. The magnetism at the end which now towards the pole, becomes the opposite of the pole was approached. If drawn from end to end in the same in rection over the other pole, the magnetism will be coincided, if not destroyed or reversed; if contrarily, it will be confirmed, if not strengthened. Hence, there are two ways of producing the same results, drawing similarly are the same pole, or dissimilarly over different poles.

Pig. 39.

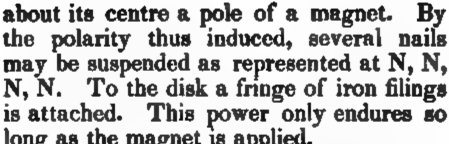
or sieve, they will arrange that selves as represented in the selves.

joining figure. Each particle of the filings become a magnet, having of course a south and north pole. The magnets thus created, arrange themselves in files, in consequence of the attraction between their oppositely may netized extremities.



731. In the files thus formed by the inductive influence of the north pole the U magnet, each little magnet will have its south pole the nearest, while in the file formed by the south pole of the U magnet the files nearest the latter, will have the north poles nearest. Hence, the extremends of the files acquire opposite polarite which are at the same time the opposit of those of the poles, with which they are respectively associated, and hence the extreme ends converge towards each other when sufficiently near.

732. Figure 40, serves to represent the magnetic power, given either to a disk of to a star of sheet iron, by applying to it:



long as the magnet is applied.

733. If a prism of iron be attracted by a permanent magnet at either pole, the north pole for instance, the prism is converted temporarily into a magnet, acquiring at the place of contact a polarity the opposite of that of the pole to which it is presented. In other words, it acquires a south polarity, when it is presented to a north pole, a north polarity when presented to a south pole. Meanwhile, the other end of

he prism acquires the same polarity as the pole by which he charge is effected. Hence, when in contact with two qually powerful bar magnets, as represented in figure 42, . prism of soft iron will be attracted with more force than

he sum of their separate forces.

H

734. Further, if after having attached a prism of iron to the north pole of one bar magnet, and the south pole of the other, on bringing the two poles remaining free into contact with each other, or with a second prism like that above mentioned, the attractive power of the four poles will be more than twice as great as that exercised by either pair\* when the others are without due communication.

735. Hence the superiority of the power of one duly magnetized bar in the shape of the letter U, ig. 41, over two bar magnets. A magnet, thus formed, is alled usually a horse-shoe magnet, from its having some esemblance in shape to a horse shoe: but preferably on count of its greater resemblance to the vowel U, it has of late been called a U magnet, by Dr. Page, and others.



 Fig. 42, will convey a correct idea. of two bar magnets as they are advan-tageously kept in a box with two arma-tures usually called keepers, on account of their sustaining the magnetic power when kept in content with the content when kept in contact with the poles as represented.

of soft iron attached to both pupon them inductively, so as a heighten their power, since it is appended to it is gradually incacquired to a certain point. O always accompanies a U magnarmature. Some keepers have tachment of weights.

N S A N

737. Thus the keeper or arma shoe magnet to attached. The be perceived, co so combined, with the same side. Thus a magnet to hold weight.

738. Under 1

cal induction, it will be seen the means of affording a conpatients to electrical discharge character, but of which, the exembles the shock given by small pairs.

#### REACTION OF MAGNETS WITH

739. The student has been deviations of a magnetic needl fluence of an electrical curren was expedient to make a prem of detecting the currents resu mo-electric influence (306).

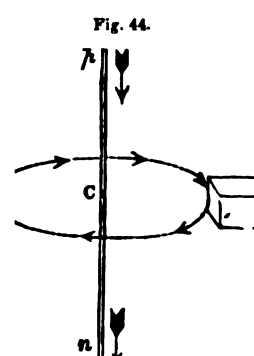
740. In order to form an ide to an electrical current which a sume could it be free from th gravitation and magnetism, let cularly on the pivot of his wat row representing the needle, as north pole points, to be attachhands, and at right angles to i



which the motion usually takes place in clocks and teches. If a galvanic current be supposed to flow through wire passing from the student's eye, through the pivot, situations of the arrow, during one revolution of the nd, will give him an idea of all the positions which the edle, when carried round the wire, would assume, relaely to the galvanic current.

741. Supposing the current to be reversed, to flow toirds the eye, it is only necessary to imagine the moveent of the hand reversed, and the arrow head pointing in
e opposite direction, to give an idea of the positions, and
e directions, which the needle would then be found to

sume if carried round the galvanized wire.



742. Suppose p, n, fig. 44, to be the axis of a circle C, and that a current is flowing through the axis in the direction of the arrows. The arrow heads in the circumference show the direction in which as many magnetic needles would point if exposed to no influence besides that of the current. When the current flows from N to P, the relative position of all the needles will be the reverse of those represented.

743. It is not possible to contrive any apparatus, which ll illustrate the reactions of a current and needle, exceptg the three instruments known severally as the compass, e astatic, or the dipping needle. These serve to illuste the phenomena only when they take place in an horintal or a vertical position. Yet from the results obtained these, an idea may be formed of those which would ensue any other positions in which needles could be situated. 744. Figure 35, already given, represents the needle in e usual form. But the directive influence of the earth is oided by associating two needles as in the subjoined figes, so that there is at either end a pole of each kind. If ese be of equal energy, when thus combined reciprocal sutralization ensues, as respects their reaction with terstrial magnetism. Such a needle is called astatic. When galvanic current flows either above or below an astatic

<sup>\*</sup> See engravings and descriptions of galvanometers.

needle, excepting that as one what nearer to the current than them will be neutralized no 1

Pig 45.

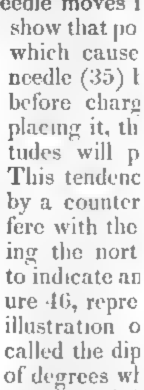
rent is ma tatic pair, i rates with a 745. Fig needle, in

ence of the

suspended, south pole a means, if the erful, the in is neutralize on the othe

746. Figure

needle in ano two slender U
747. In the ters needles ar hair or filamen cocoon. By the earth over lized. In anoth needle moves i



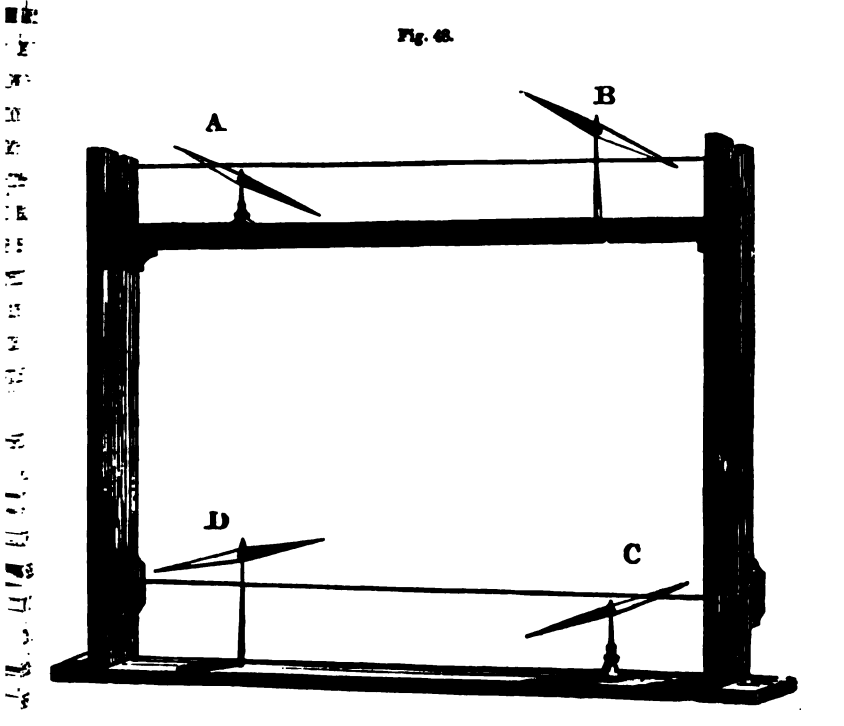
zontal level, i



Fig. 47



748. The following engraving represents an apparatus, means of which, four needles A, B, C, D, are simulta-



neously exposed to the influence of the same current, in respect to which they occupy different situations. Of any two needles, of the four, one of which is above the other below the current moving the same way, the poles will be seen discordantly directed. Of course any two of the four of which one is above the current moving one way, and the other below it, while moving the other way, will agree.

749. Accordance in the direction of needles exposed to the same circuit, is found only when the situation relatively to the current, and the direction of the current are alike, or where diversity in one respect is compensated by diversity in the other. In order that the currents thus moving in opposite directions should not counteract each other, it is necessary that two or three feet should intervene between those portions of the wire in which they take place.

750. The apparatus must, in order to show the phenomenon, have the wire in the plane of the meridian, so as to have the needles parallel to it; as, when uninfluenced

71

2

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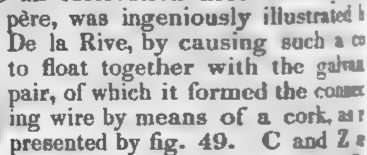
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by the current, they place themselves so as to commit with the meridian plane nearly, or, in other words, we

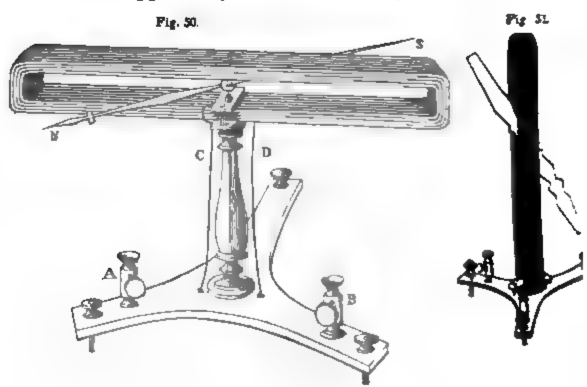
to point north and south.

751. The necessity which exists, of performing the appearance to as to avoid every injurious influence from the restrial magnetism, ought not to create the idea that in phenomenon is dependent upon that magnetism. We astatic needles (44, 45), in which the polarity of cocard die is so counteracted by that of the other, as to rest the two indifferent to the polarity of the earth, the dentitions may be produced without reference to the mendature.

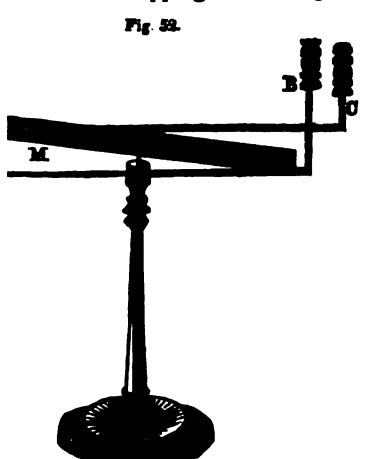
752. The magnetism imparted to a coil by a galuar current, agreeably to an observation first made by An



small plates of copper and zinc, above which the coll supported, the ends being soldered severally to the plate. When the whole is made to float upon acidulated water galvanic current passes through the coil from the copper to the zinc plate. Under these circumstances the codure react like a magnet with either of the poles of a magneticiently approximated, so as to cause the whole apparatus to be apparently attracted or repelled.



. Two forms of the galvanometer (50, 51), are illustrated by the prefigures, one of which represents the situation of the needle when rted in a horizontal, the other in a vertical plane. Figure 51 corids with the dipping needle; figure 50 with the compass needle.



754. The rationale of these instruments may be made evident by the more simple form represented by figure 52. In this the wire makes but one circumvolution, as in the larger apparatus, fig. 49. The influence of the simple current being understood, it will be easy to conceive, that if the needle were subjected to two such circuits, the effect upon it would be arithmetically greater as the number of such circuits acting upon it from an equally favourable position. Consistently, Schweiger, by multiplying the circuits or circumvolutions, produced the famous instrument in question, appropriately called a

lier as respects its origination, but not as to a designation of the use

ch it is applicable.

. In multiplying the circumvolutions, it is not necessary to multiply teries by which they are supplied, since, when formed of a continuous he first and the last may, from the same battery, be supplied, if not y at the same time, with an interval almost infinitely small. each circumvolution to measure a foot in circumference, twentyndred would be only half a mile, and this distance would be gone h in less than the five hundred thousandth part of a second, agree-, the inferences of Wheatstone, supposing them applicable to currents intensity, as well as those of the statical kind.\* For a galvanomenstrument of the highest sensibility, see Melloni's thermo-multiplier, the head of Thermo-electricity.

enometer or Multiplier of an unusually large size, made of Strips of Tin wound in a Coil with interposed Strips of very thin Paper.

engraving represents a large multiplier, or galvanometer, the needles of are each about eighteen inches in length. The instrument is furnished above circle graduated into 360 degrees. Agreeably to the usual construction, the being within the coil, is subjected both above and below to the concurring ce of a current passed through the coil. Under these circumstances, the peodle is situated in the apparatus represented by the adjoining figure. In nation which the upper needle occupies, the influence of the lower portion of l, so far as it operates, must counteract that of the upper one. Yet when the portion of the metallic coil is at a distance from the upper portion of about rd of the length of the needle, and this is situated very near to the upper porhere represented, the influence of the latter may so far predominate as to the indications very nice; while they are much more easily seen and estiby means of the graduated circle, when, as in the situation of the upper nothing intervenes between it and the eye.

nother instrument of the same dimensions, I have used only a semicircle for

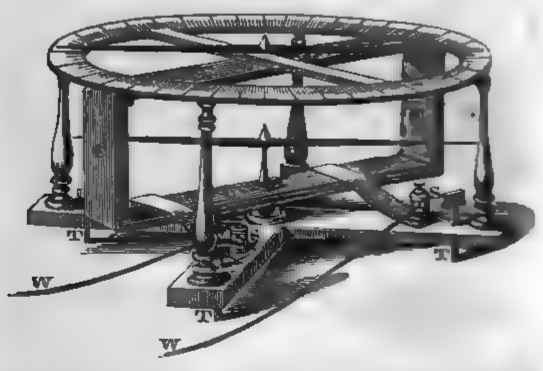
duations, which, excepting the appearance, answers as well.

ou of wire, a coil of tin foil of about an inch in breadth, and eighty feet in

#### Ampère's Revolving Battery.

756. The most simple process for creating a rotary motion, by of the reaction between an electrical current and a magnet, is that is with





length, separated by thin paper, may be used, but a copper wire of No. 16. 25 about 180 feet in length, coated with shell lac varnish, will be more efficient. The coil of tin-foil or varnished copper wire, is wound about the paralleless. C, C, C The ends of the coil are severally soldered, or screwed, under the la

the gallows screws, S, S

When both needles are placed upon the pivot at the same time by the sep of their similar poles, they will diverge from the meridian unloss they be versed situation, in which case they will both appear as in the engraving. pole of one needle pointing north, the north pole of the other, so ith these circumstances, a discharge is made through the surrounding cold. De-

The clean surfaces of disks of zinc and copper, each an inch in diameter. by paper moistened with pure water, are sufficient to move the needles =: The wires, W, W, are used for the purpose of connecting the disks with the? the wires. They are attached to the instrument by gallows screws, S. S.

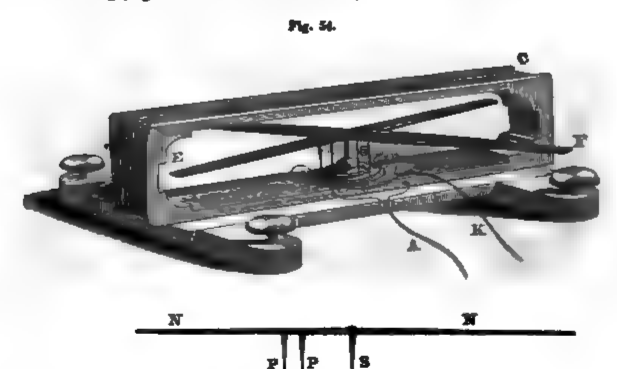
The level of the machine is preserved by the aid of four screws, of which three can be seen in the drawing at T, T, T.

Rotary Movements arising from the Reaction between Magnets and Ga. Currents

ENGRAVING AND DESCRIPTION OF A ROTARY MULTIPLE Or one in which one or more Needles are made to revolve by a Galranie C

The preceding engraving represents a rotary galvanometer, or multiplier a contrived in November, 1836, and which must have value as an add.com amusing, if not to the useful implements of science. It is well known t passing a temporary discharge through the coil of a multiplier, the needer made to perform a revolution, whereas if the current be continuously apply movement is checked as soon as the situation of the poles is reversed. To p a permanent motion, the discharge must be allowed to take place only wh poles are in a favourable position relatively to the excited coil. This obje tained by means of two pins, descending from the needle perpendicularly a enter two globules of mercury, communicating, on one side, with a galvanic; the other with the coil of the multiplier. In the next place, by winding or first coil, another of similar length, but in a direction the opposite of that is

**the** galvanic triad, by which the current is generated, is made to revolve about the pole of a magnet, which serves as a support. This is exemplified by what has been called Ampère's rotating battery, of which figure 55, on the following page, will afford a satisfactory idea.

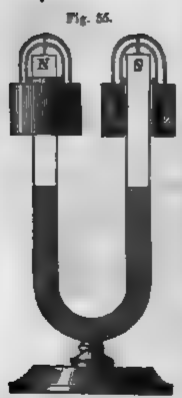


the first coil was wound, I was enabled, by two other globules, situated so as to communicate severally with the lower ends of the pins, at the opposite side from that on which the first mentioned globules were, to cause an impulse at every semi-revolution.

The one coil being wound to the right, the other to the left, the alternate effect of each upon the needle was similar in opposite parts of the orbits described by the pine. Lastly, a second needle, furnished with pine in like manner, being fastened at right angles to the first, so as to form with it a cross, as represented in the engraving, each needle is made to receive two impulses during every revolution. Hence one of Daniell's sustaining betteries, as made by Newman, is quite adequate to cause a revolution as rapid as consistent with a due degree of stability in the mercurial globules employed.

One end of each coil, by means of the branching wire A, communicates with one pole of the galvanic pair; the other ends of the coils terminate in mercurial globules contained in cavities on opposite sides of the wooden disc G, upon the centre of which the spindle of the magnetic needle rests. The branches of the wire K proceeding from the other galvanic pole, terminate in globules situated in the vicinity of those above mentioned, so that as the needles revolve, the pins proceeding therefrom perpendicularly may touch a pair of the globules first on one side and then on the other. Whenever this contact takes place, the circuit is completed, and a discharge is effected through one or the other of the coils of the multiplier.

Bupposing E and F to be north poles, a discharge through one of the coils will cause E to move off a quarter of a circle, or more. As this ensues, the pins of F will come in contact with the globules which those of E touched before. Of course F will be propelled so as to cause the pins of E to reach the pair of globules at G, which, completing the circuit of a coil wound in a way the opposite of that first mentioned, concurs with that coil in its influence, so as to promote the rotation previously induced. The same result ensues when the pins proceeding from F, come in contact with the globules situated at G, and when E returns to its original starting point. It follows that by a repetition of the process the galvanic action is sustained. The phenomenon is as well illustrated by employing the single needle, N, N, as by two, but the most pleasing and energetic effect is produced by the crossed needles. In this simple form the spindle on which the needle rests and revolves is represented at S; the pins at P, P. Each coil, consisting of copper bell wire, is about thirty feet in length, and is contained in the groove C. The frame of the multiplier is constructed of mehogany, and is levelled by the milled headed screws, on the ends of which it is supported.



757. Two concentric cylinders of sheet case: one about three-eighths of an inch larger on the other, are united by a flat ring of the same care. so that the latter forms a bottom, while there are tween them an interstice of three-eighths of in ac-The inner cylinder is open and furnished with arch of copper, at the summit of which then at pivot pointing downwards. This pivot rests up a agate cup seated upon one of the poles of a semagnet. By means of another arch of case: cylinder of zinc is supported within the interests. resting by means of an appropriate pivol co a 😂 concavity in the lower arch above described, as == porting the copper cylinders. Upon the care as of the magnet another similar apparatus of colors is supported. When the apparatus is to be put =: operation, the interstices between the copper cyaders must be supplied with a diluted mutan 4 nitric and sulphuric acid-A galvanic current lat ut place in each little battery through the supported arches from the copper to the zinc (320), the cas-

the two batteries, the similar cylinders move oppositely. In

dissimilar cylinders move alike.

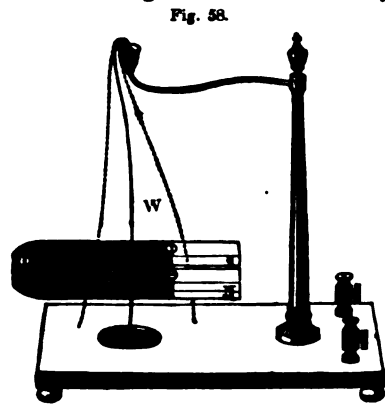


758. A magnet, n s, being attached, as in figure 16, by a silk thread, to the centre in the bottom of a conf mercury, so as to float in that metallic liquid, a wet. 4 b, is fixed over it vertically, so as just to enter the arrcury for a small depth. The metallic support of the wire is connected with one pole of a galvanic better. the mercury with another. In this case, the utner roo of the magnet, while swimming in the mercary, my about the wire from right to left, or from left . ... accordingly as the poles of the magnet, or the a magnet, tions with the galvanic apparatus, may be variety on the other hand, the magnet be made in to ar while the vertical wire is upon a universal year. wire will revolve about the magnet, and be to change its direction, in the same way, as " moveable magnet was actuated, with respect to to

759. The letters n and s, and direction of the atrows, in either of the adjoining figures, representing the apparatus alluded to, show the relative situation of the poles of the magnets, and direction of the current requisite to produce a revolution from right to left. In either apparatus a reversal of the relative position of the poles, or of the direction of the currents, will reverse the rotary motion. Of course n designates the north, and s the south pole.

760. When one end of a wire is fastened to the axis of the pole of a magnet, leaving the other end free, it will revolve about the point of attachment as a centre of motion, and in like manner, if the magnet is fastened

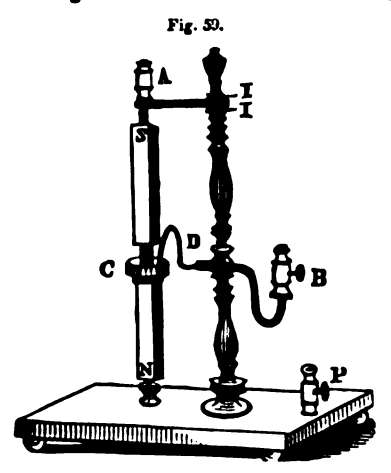
The wire, because in either of the cases, whether it be that of the wire or magnet, an escape from one point of collision existing in the circle, throughout which the conflicting influence prevails, only carries it to another such point. Thus the point of attachment, about which the motion takes place, becomes a centre, about which either the magnet or the wire revolves, while making Ixion like efforts to fly from inevitable collision.\*



761. Let the end of the wire W hanging from a metallic bracket, which communicates with one of the set screws on the right of the platform, enter some mercury, occupying a cavity in the latter, which has a metallic communication with the other set screws. Under these circumstances, as soon as the set screws are connected with a galvanic battery in operation, the end of the wire flies out of the mercury, but the circuit being thus broken, the end of the wire soon falls again into contact with the mercury, and from the same cause as at

" I here quote from Daniell's Manual a very interesting illustration of the reaction of a galvanic current with a bar magnet.

"Magnet revolving round its own Axis.—The instrument represented in figure 59 is designed to show that the action between the current and the magnet takes place



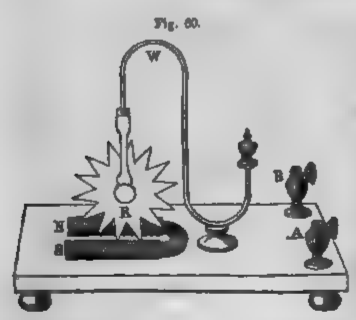
equally well when the magnet itself forms the conductor of the electricity. The lower end, N, of the magnet, being pointed, is supported on an agate at the bottom of a brass cup connected under the baseboard with the binding screw cup, P. The upper end, S, is hollowed out to receive the end of the wire fixed to the cup, A; the brass arm supporting this cup is insulated from the brass pillar at I, I, by some non-conductor of electricity. To the middle of the magnet is fixed a small ivory cistern, C, for containing mercury, into which dips the end of the wire, D. Thus the magnet is supported with its north pole downwards, and is free to rotate round its vertical axis. A little mercury should be put into the cavity at S, and into the brass cup at N, and the ivory cistern be filled sufficiently to establish a connexion between the magnet and the wire, D.

"On connecting the cupe, A and B, with the battery, the current will flow

through the upper half of the magnet, causing it to rotate rapidly. If the cups, B and P, form the connexion, the current will traverse the lower half, equally producing revolution of the magnet. Now connect A and P with the battery, and no motion will result, because the electricity passes through the whole length of the magnet in such a manner, that the tendency of one pole to rotate is counteracted by that of the other to move in the opposite direction. Connect B with one pole of the battery, and A and P both with the other pole. The magnet will now revolve, since the current will ascend in one-half of its length, and descend in the other."

first, again flice out. Thus an alternate motion is produced and mans as long as the current is continued.

#### Barlow's Revolving Wheel.



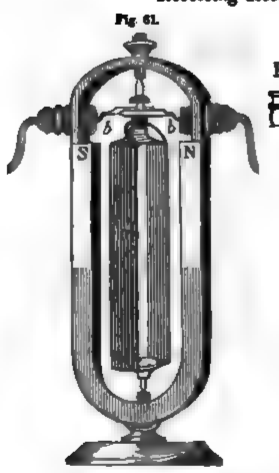
762. The apparatu == sented by figure 60, a see Barlow's revolving wim, the name of the invests b wheel being situated being the jaws, either of a star at net or an electro-mand w made the medium of the pr vanic current, say toth # contact with the mercay = > cavity under it, is across to the same way as the sate the experiment above durable (762). But when agreesly the same law, one tout in from collision with the most

Every tooth being successively affected in like manner, the where was long as the current endures. It is not necessary to the motion who circumference be serrated, but it serves to make a pretty spark and contact. The endurance of light thus arising, is too brief for the whole make a perceptible movement while it lasts. Hence, when the natural is illuminated only by its own scintillations, it appears to be at rest, because rapidly it may revolve. It is computed that eight sparks in a second protect of a continuous illumination, and as the wheel gives now these, it appears as if seen by a continuous light.

Explanation of the Motions produced by means of the Apparatus scribed in the three preceding pages.

needle free to move, and a wire transmitting a galvanic current, that the former will strive to assume a position at right angles to a railed to the axis of the wire. Of course, as action and reach a result tionary needle, or other magnet, will arrange itself so as to product same relative position between the two bodies as when the wire is save ary, and the needle free. But if freedom of motion exist only a constitution, whether of the needle or the wire, the motion can only take at that extremity. Yet if the movement hence arising be not such as the terrupt the galvanic current, or be such as to permit its renewal as some discontinued, the consequent movement must be continued or reposition may be productive of rotary motion, as in the case of figures 54.55.56, 59, or of alternate motion, as in the instance of 58.

#### Revolving Rectangular Coil.



764. As in the galvanometer of Schweiger, the force of the conflict between the dynamic polarity of the current, and the stationary polarity of the needle, is multiplied by reiterating the presence of the former in the coiling of wire through which it has to pass, in order to perform its circuit, so the instrument of which figure 61 is a representation, is an apparatus in which a current transmitted through the rectangular coil, between the jaws of the magnet, is made to react with the magnetic influence of the latter, and strives to escape therefrom, in order to assume the position in which there will be least in-This takes place when terference. the plane of the coil is so situated as to be at right angles to a line passing through the centre of each pole.

65. In fact the coil becomes a magnet, (752,) and of course tends to age itself, as would a magnetic needle similarly situated; but, by means a contrivance called the pole changer, described below, in making a i-revolution to effect this arrangement, the direction in which the curgoes through it is reversed, and consequently a different impulse is sived, causing another semi-revolution. No sooner is this made than impulse again changes. Thus a continuous revolution is produced as a sthe current endures.

#### Of Page's Pole Changer.

66. In some cases, in order to produce opposite impulses, it is necessary sause a galvanic current to flow alternately in opposite directions, in ers, as it will hereafter be seen, it is an object to render an alternating rent constant in its direction. For these purposes the following contrice was devised by our countryman, Doctor Page, and executed by Daniel ris, jr., the highly ingenious artist, to whom we owe the successful fabrion of a variety of electric, galvanic, and electro-magnetic apparatus.

67. The apparatus in question being employed to reverse the polarity the electro-magnet in the apparatus last described (767), the following



explanation is given in reference to the use of it therein made. The pole changer, figure 62, consists of two small segments of silver S, S, fixed on opposite sides of an axis of which a section is represented in the figure. These pieces are not in contact, and are sufficiently insulated from each other, and from the

s. Of the two ends of the coil, situated between the legs of the magnet represented, one is soldered to each segment. By means of a wooden h, of which the polar ends S, N, of a U magnet are the abutments, two

wire springs, b, b, are so supported and insulated, that through the mane tion either of two cups of mercury, or two set screws, at B, B, throme be made to communicate at pleasure with a galvanic battery. The impact of these wires are made to press against the argments of a most sufficient force to make a good contact. The arrangement is such the respective contacts of the semi-cylinders with the springs will the parameters when the coil C is, as represented in the figure, at right angles to a parameter of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will the parameters are the coil C is, as represented in the figure, at right angles to a parameter of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will the parameters of the semi-cylinders with the springs will be parameters of the semi-cylinders with the springs will be presented in the figure, at right angles to a parameter of the semi-cylinders with the springs will be parameters of the semi-cylinders with the springs will be parameters of the semi-cylinders will be parameters.

passing through the axes of both legs of the magnet.

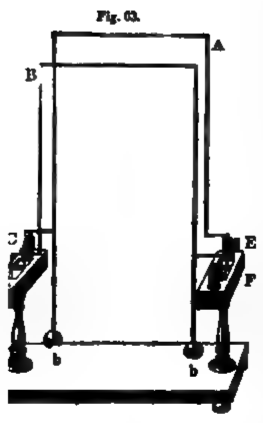
768. The apparatus being thus prepared, let the mercury copied course the wire springs, be in communication with a competent bater, as sequently the current will from one semi-cylinder, enter one end of a coil, proceed through the circumvolutions of the latter, and go out a cother end of the coil to the other semi-cylinder; but if the con to the half round, the segments, and the ends of the coil attached to them man tively, will have their electrical states simultaneously reversed, create the sides of the coil a corresponding change of electro-magnetic policy. Thus when the coil is turned from the position into which the position would bring it, in seeking to regain that attitude, it reverses them. The causes it to make another half revolution, which equally operates to reconstitute propensity to change, and thus rotary motion is induced as long at a current is sustained.

# Attraction and Repulsion between Electrical Currents of the Movements thence arising.

tained by Mr. Ampère, a French philosopher of celebity to whose inventions and theoretic suggestions, electromagnetism is pre-eminently indebted, that wires, completing the circuit of different galvanic batteries, would attract and repel each other; but not in a mode analogous to electrical reaction, since it was between the bodies, similarly excited, that the attraction was observed, while the represent took place between bodies dissimilarly excited. Electrical indications are obtained only, when the poles of the generating apparatus are unconnected, those afforded by the galvanized wires, were the consequence of their connecting the poles of the galvanic apparatus.

770. Figure 63, represents an apparatus, contrived for the purpost a shewing the reaction between wires subjected to galvanic currents.

771. Two copper wires, A, B, are supported parallel to each other, a such manner as that their upper parts may move freely towards each the so as to touch, or in the opposite direction so as to become more recording ties replete with mercury, being amalgamated so as to insure a project contact with that metal. By means of the little metallic balls, b, b, which mercury be lowered or raised by screwing, the centre of gravity of the wires my be so adjusted as to keep them perpendicular, when undisturbed, and we



render it easy for them to oscillate from a very slight impulse. The four cavities holding mercury within which the points rest, as above stated, have each a metallic communication with the set screws C, D, E, F, when E, F, are made to receive wires from the same pole of a battery, while the other two, C, D, communicating with the other pole, the current will go the same way through both of the wires. Under these circumstances they will attract each other, and come together; but if E and C, be made to communicate with the one pole, while D and F, communicate with the other, the current will flow oppositely in the wires, and they will move apart, as if actuated by repulsion. If the wires be made to oscillate by a timely opening and of closing the circuit, the motion will be increased.

tion of one Coil of Wire within another, in consequence of the rapid eation and reversal of the Polarity arising from the change of direcm in a Galvanic Current caused by Page's Pole Changer.

Pig 64.

772. The adjoining figure 64, is designed to illustrate by means both striking and agreeable, the powerful and rapid attraction, and repulsion, between wires caused by electrical discharges made through them alternately in opposite directions.

773. This apparatus has a close analogy with that already described figure 61, as to the means by which the polarity of the coil is imparted, and the direction reversed at every semi-revolution, by means of a galvanic current, and the pole changer (767). We have only to substitute for the U magnet and rectangular coil, two circular coils of the same material as the rectangular coil. Of the two coils, one is large enough to contain the other without touching, and being stationary, takes the place of the U magnet. The other,

oting shape, does not differ from the rectangular coil; while it is persimilar, as respects the means by which it is supported on an axis
associated with the semi-cylinders of the pole changer, and the office
h it performs. In one case by polarities arising from a current, the
ugular coil is made to rotate by conflicting with the poles of a U magin the other case the polarities arising in a circular coil from the same
t, conflict with those arising from a current in another coil. In either
seeking a non-conflicting attitude, it is disappointed by the consequences
own efforts, and the perverting influence of the pole-changing appa-

## Of Electro-dynamic Induction.

1st. Under this head we may put Induction of a Current upon it 2d. Generation of Currents by the Dynamic Inductive Information Currents.

Induction of a Current upon itself.



774. Figure 65 represents as of copper, in weight thirteen pr in width one and an half ind length ninety-three feet. This is wound into a flat coil, of a inches in diameter, but, being im by a double coating of sile, that be no metallic contact between of its circumvolutionsend of the coil thus formed, in. presented, connected with one an active battery e, to the our of which a rasp d, is soldist scraping the rasp with the free the coil, sparks and metallic da tions ensue with much more con scintillation and louder snapping

than if the same scraping were to be performed with the terminate conductor having a length no more than requisite to complete the to Moreover, Henry, I believe, first observed, that an operator touchi rasp with one hand while holding the end of the coil during the set would be subjected to a perceptible shock at each of the interruptions circuit, to which the sparks were indebted for their existence.

ceding paragraph, whenever the ribbon touches the rasp, the late two circuits at command, one completed altogether by the coiled ribbon touches in part by the person of the operator. Yet on account of the itensity of the current, the imperfectly conducting circuit formed by it mal frame, will not be the mean of a perceptible discharge so long other route afforded by the perfectly conducting copper ribbon is a rupted: but when the ribbon circuit is ruptured, a perceptible distance place through that made up of the operator's person, the coll he holds, and the rasp which he touches.

petent to produce any perceptible discharge through the operator. I quently the shock, produced as described, and all the other character in which the phenomena arising from the breach of the circuit where coil is interposed, differ from those which appear when a short metalliductor is similarly employed, must be ascribed to the intervention coil. It would seem, therefore, that the materials of the coil, both poble and imponderable, become coerced by the battery into an uncestate; so that, on the rupture of the circuit to which this state is distant recoils towards its natural equilibrium, producing an electionarge of an intensity far above that of the parent stream sent through coil from the battery. On this account, it would seem as if the redischarge might be appropriately designated as a recoil current. It conductor, which is the channel of a galvanic current, is in an ar

ism seems to justify. This state, noticed by Faraday, was designated by him as an electrotonic. The tension arising from this state, or the energy of the effort to have it made by the matter in which it has been produced, appears to be greater when the conductor, being in the ribbon form, is wound up as a coil. Doubtless the proximity of the circumvolutions to each other, must cause a reaction between the similar polarities, tending to augment their resistance to the state imposed on them by the battery.

within the ribbon coil, is that which arises in a voltaic series of sufficient intensity when the circuit is incomplete. Hence, agreeably to Henry's observations, the direct action of a series of high intensity, forming a circuit through a short conductor by sudden contact, resembles, as to intensity, the indirect action of a battery of low intensity, on breaking the circuit formed through a long conductor. In the one case, the intensity being due to the resiliency of the electrifiable particles of the conductor, is increased with its length to a certain point; in the other case, the intensity being due to the battery, the conductor cannot be too short: excepting that when the effect is measured by the current inductively created in another conductor, agreeably to the process next to be explained, this effect must be influenced by the length of the conductor, as well as the intensity of the current which it conveys.

778. It has been suggested (776), that the electrical discharge accompanied by a shock and unusually large and noisy spark which characterizes the process under consideration, is the consequence of the sudden effort of the matter of the coil, both ponderable and imponderable, to recover its natural equilibrium from a state of forced polarization within the coil. There is a rough analogy between this resiliency of the supposed imponderable current, and that which would ensue, if, instead of a copper coil dynamically electrified, a steel watch spring were wound up, to a certain degree of tension, and then suddenly liberated so as to be allowed to react.

779. But in the act of recoiling, if, by the discontinuance of the broad channel in which it may be generated, the current be diverted into a collateral channel, in which it must experience far more resistance, the animal frame of the operator, in the case in point, the concussion must be proportionable to the impediments, just as if a torrent by being suddenly dammed up, were forced into a circuitous channel, containing rocky obstructions.

780. In the instance of a principle, which moves with the inconceivable velocity which characterizes electrical discharges, the larger the mass which recoils, the greater the effect; but as to wind up a spring instantaneously, requires an activity in proportion to its length, the longer the wire in which the tension is to be induced, the greater the voltaic intensity required. Thus Henry found, that when a voltaic series, consisting of six pieces of bell wire, each one inch and an half in length, and an equal number of pieces of zinc, of like dimensions, were employed to excite a due electrical tension in a spool of wire five miles long, and one-sixteenth of an inch thick, the shock resulting from resiliency on the rupture of the circuit, was sufficient to affect twenty-six persons joining hands.\*

I quote here from Davis' Manual of Magnetism, an engraving and description of an instrument designated as the Contact Breaker. Supposing to employ this mechanism in the production of the phenomena, arising from a rapid rupture and remewal of a galvanic circuit, this explanation may be desirable to some of those who may witness the processes in which it may be used.

# Generation of Currents by the Dynamic Inductive Informat of the Currents.

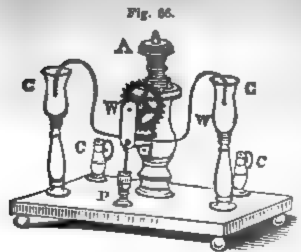
781. The power which an electrified body has to produce an open charge in another, has been explained under the name of "Electron's citement by Induction" (E, 107). There is another species of second influence, to which allusion was made as dynamic induction (E, 120).

782. The attractive and repulsive influence of wires subjected to proceed that there is such a reaction between the cause of electricity in transcent conductor, with the same principle in another, that under favorate as cumstances a current in one may inductively produce a transient current the other. Thus, when two long wires were associated in concerns the other. Thus, when two long wires were associated in concerns the with which the wires had been covered like bonnet wire, on makes was produced through the other. This effect was detected by subject to needle to the dynamic discharge through a small hollow helix of with cluded in the resiliating circuit.

783. The following laws have resulted from Faraday's researcher.

"1. During the time a galvanic current is increasing in quantity and ductor, it induces, or tends to induce, a current in an adjoining parallel.

It consists of a bent copper wire W W, which by means of clock and motion by a spring, is made to vibrate rapidly, dipping its ends alternated glass cups G G, intended to contain mercury. The spring is wound up by the milled head A. The glass cups



the milled head A. The glass cope open at the bottom to allow the acceptance of the come in contact with the brass plant which they are comented. They are both connected with one of the large both connected with one of the large both connected with one of the large both connected with one of the municates with a brass mercury and into which dips a short wire committed which dips a short wire committed with the vibrating wire. S. L. 22 are carry must be put into the capt. I will could be end of the vertical with the capt. I will could be completely the before the other can be its portion.

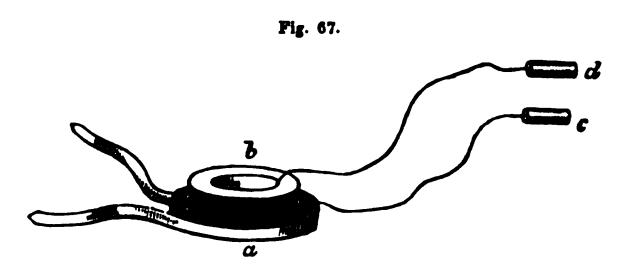
The Contact Breaker may be 1.75

of the instruments for affording sparks and shocks, which will be described the following head. The current must be transmitted through the two lessmoods in succession, by connecting one of the cups C.C. with one pole of the bullet the other cup with one of those attached to the spiral or other piece of apparent the remaining cup of which is to communicate with the other pole of the better is better to break the circuit mechanically in this way, than by means of any sprupting apparatus worked by the battery itself, as a considerable part of the piece of the current is then expended in giving motion to the interruptor.

On making connection in this manner with a flat spiral, and turning the me head A to put the vibrating wire in motion, a brilliant spark will be seen it were rupture of contact, accompanied by a loud snap, and producing considerance bustion of the mercury. With a battery consisting of a few pairs of plates. Are size, such as Dr. Hare's Calorimotor, the size of the spark will be greatly in the and the snap become as loud as the report of a Leyden jar. The shock will as pretty strong, and may be increased by covering the mercury in the glass cups a stratum of oil. A shock may be obtained, especially when oil is used on cover the circuit as well as on opening it, though inferior to that given in the litter as a faint spark is also sometimes seen when the wire dips into the increasy

rimary current in full quantity, no inductive action is exerted. 3. But rhen the same current begins to decline in quantity, and during the whole inne of its diminishing, an induced current is produced in an opposite direction to the induced current at the beginning of the primary current."

784. Professor Henry, of Princeton, has shown that currents may be trikingly produced by induction, in the following manner. The circuit bruned, through the ribbon coil, a, with the battery, figure 65, being alterately made and interrupted, as stated (774), a helix of coated copper wire about 21 wire gauge, and of about 4900 feet in length, is subjected to the nductive influence of the coil, as represented in the adjoining figure, in



which the coils and helix are situated concentrically one upon the other, while separated by a plate of glass. The intensity of the resiliating current as measured by the shock, increases with the length of the helix employed, until the resistance to the electrical current consequent to the length, compensates the benefit otherwise arising. Yet, as might be expected, the length to which the circuit may be thus extended with advantage, ceteris paribus, is proportional directly to the numerical extent of the voltaic series employed (431).

785. The ends of two wires severally connected with those of the helix, being rubbed one against the other, while the circuit through the coil is alternately broken and renewed in rapid succession, sparks will be visible, and slight shocks may be felt through the fingers or tongue, on contact with the wires. When the ends of the wires are joined, the sparks and snaps, which can be obtained between the end of the coil connected with the battery while passing over the rasp, are much diminished in energy, and no shock can be received.

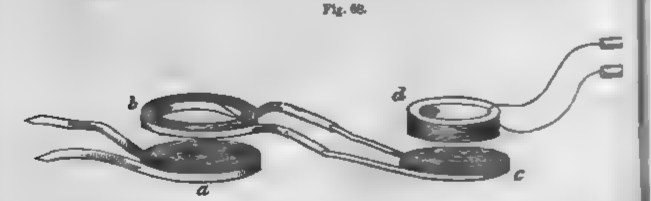
786. A delicate galvanometer being associated in the circuit with the upper coil, whenever the battery circuit is completed through the coil, the magnetic needle will be extensively deflected, but will immediately return to the meridian, a flow of a current through the wire of the galvanometer is thus indicated, which is nevertheless transient apparently. On opening the circuit, a similar transient deflection will occur in the opposite direction. No deflection occurs while the battery current is flowing steadily. The galvanometer should be placed at such a distance from the coil that the current through this may not affect the needle.

787. A sewing needle will be magnetized if placed within a spiral of wire of very small diameter, duly comprised in the same circuit as the belix. The polarity produced by the current which ensues on the completion of the circuit, will be the reverse of that communicated by the current consequent to its rupture. If both currents are allowed to act on the

peedle, it will acquire little or no magnetism, as the magnetizing misses

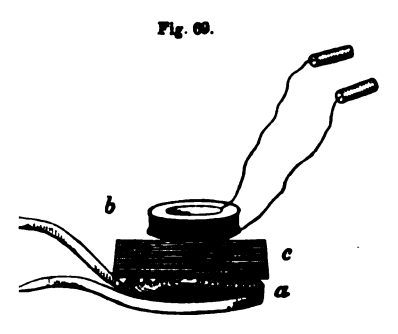
of one will counteract that of the other.

788. Instructed by Faraday, Prof. Henry has added many inguises acoveries respecting the phenomena of dynamic induction; among these the consistent consequence, that the transient accondary current at the described, inductively produced in a second coil, was capable of text remainded conductively, so as to circulate through a third coil, when are ing inductively on a belix, was productive of a tertiary current of the second character, though feebler, as that produced when the limits is expend to be inductive influence of the primary coil. Hence shocks will be recently



a person holding in each hand one of the handles, to which the ends of the helix are attached, as represented in figure 68. The process may be tended so as to produce, analogously, fourth and fifth currents. "Heart alleges that a shock was given, by means of a current of the third order, the twenty persons joining hands." Shocks extending to the arms were the produced by the process above mentioned, "with currents of the felt of der."

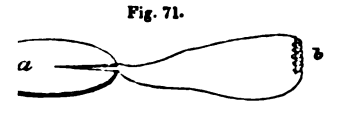
789. Of course, the characteristic efficacy of the primary current and be liable to vary with those of the galvanic apparatus employed a 🖘 🗯 duction, and the length, width, and thickness of the ribbon empers. With a simple elementary battery of a given size, there must be a comlength, width, and thickness of ribbon, which will produce a maximum of fect; and any deviation from this combination of requisites must be prosective of deterioration. It follows, that with ribbons of various discusses. there must be for each a degree of intensity and quantity depending on 🛎 number and size of the voltaic pairs, which will render the inductive power produced by its means, a maximum. Hence the sectional area and leads of the ribbon, the extent of the oxidizable surface of the zinc, and the purber of the series into which it may be divided, form variable elements. ceptible of an almost infinite variety of complications. Each of these rerious complicate combinations must have its appropriate effect upon a se condary coil, having a given length and sectional area, but must vary with every change in either of these dimensions. In like manner, the charge teristics of the current produced by the inductive influence of a second. third, a fourth, or a fifth coil, or as many as may be employed with care. must vary with their length, shape, and sectional area. Another source of variation arises from the distance of the inducing coil from that whet \* the subject of the inductive power, and also their parallelism, and the 🔄 gree in which their axes may approach or recede from coincidence was one line. The directions of the currents, inductively incited, seems have varied with the distance.



790. To Prof. Henry is due the detection of the screening power of metallic sheets. I allude here to the fact, that the interposition of a metallic sheet, as in figure 69, between the ribbon and coil, or between a coil and a helix of fine wire, nearly annihilates the power to produce shocks, although the interposition of a plate of glass, or any other nonconducting material, does not impair the inductive influence. Yet

Fig. 70.

a sheet of metal, out of which a gore or sector has been cut, as represented in figure 70, does not impair the power of induction. Moreover, when from the corners of the gash made in a plate, by cutting out the gore, wires soldered thereto were duly extended and attached to the ends of a helix, in-



cluding a needle; to this needle, on subjecting the plate to the inductive influence of a coil duly excited by a battery as in the adjoining figure, polarity was imparted, which was such as to show the resilient current to be in the

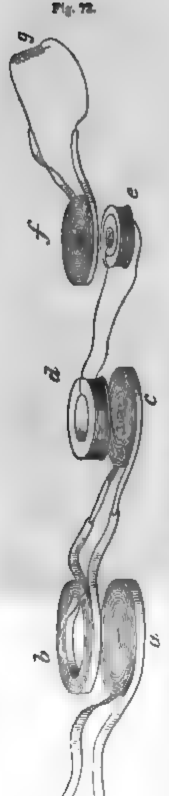
e direction as that in the inductive coil, a. This resilient current is idered by Henry as a secondary one, but it seems to me it is to the imable current of which it is the resiliating effect, that the appellation of ndary should be applied.

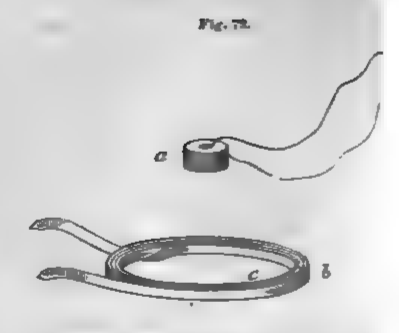
31. I find that a sheet of zinc does not prevent the creation of a resilizarrent in my helix (793), sufficient to affect the gold leaf galvanos. I am under the impression that the plate acts by causing a diffuor of the inductive force, so as to reduce its intensity too low for causing ock.

92. The screening power above described, as displayed by a sheet of il interposed between the first and second coil, equally availed, when a interposition was made between a second and a third, a third and a th, or a fourth and a fifth coil.

93. When a copper ribbon coil is made to act inductively on a long of wire, as illustrated in figure 67, the shock arising from the resili7 of the induced electrical tension seems, within an extensive boundary, e greater in proportion as the wire is longer and more slender. I have loyed a ribbon coil of about three inches and a half wide, and one hundand ninety-six feet long, and a helix of coated copper wire, No. 21, of at twenty-eight pounds in weight, and a mile in length, furnished with the metallic handles. Although the helix may be supported at the disce of several feet above the coil, while the current of a calorimotor is alately established and arrested within it, a person holding one handle of helix wire in each hand, will receive shocks which augment in severity he helix is approximated to the coil, and become intolerable when it is aght into contact with it nearly.

94. Figure 72 represents a helix such as described, situated as when it smart shocks, in consequence of the inductive resiliating influence of coil, c.





795. From the preceding statement, it epper a current of great quantity may produce a s current of great intensity. But Henry has and that the converse is possible. The adjoining fig been employed by him to illustrate the experim which the facts alluded to were established.

796. The tension induced by coil a, in on transmitted conductively to coil c, whence # 1 ductively upon the helix d, producing intensity conductively transmitted to e, whence it in generates a current of quantity therein, sun magnetize a needle included in a little spiral

as represented.\*

797. Much labour has been bestowed by H determining the directions of the current in ferent coils; but in this part of his investigation is too much abstruse and minute detail, to?; nected with striking phenomena, to be intrid. a text-book. I am under the impression, that ing out the consequences which theory would founded on the idea that each current, indoes duced, must be the opposite of that which imm induces it, would give the direction of as h generated in the coils and helices of figure 73.

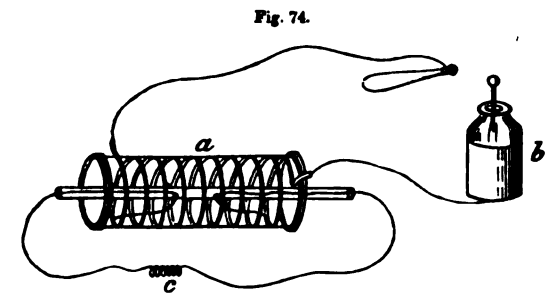
### Dynamic Inductive Influence of Statical Electricity.

798. On this subject I will quote a few lines from Henry's Methe American Philosophical Transactions, together with the appropr graving.

799. "The discovery of the fact that the secondary current, w! ists but for a moment, could induce another current of considerable gave some indication that similar effects might be produced by a di

" It must be evident, from the experiments here illustrated, that the sam tive power produces either quantity or intensity, accordingly as it is made the long and narrow circuits afforded by the wire helices, or the shorter a ample channels afforded by the ribbon coils.

ordinary electricity, provided a sufficiently perfect insulation could be obtained. To test this, a hollow glass cylinder, fig. 74, of about six inches in diameter, was prepared with a narrow ribbon of tin-foil, about thirty feet long, pasted spirally around the outside, and a similar ribbon of the same



Length pasted on the inside; so that the corresponding spires of the two were directly opposite each other. The ends of the inner spiral passed out of the cylinder through a glass tube, to prevent all direct communication between the two. When the ends of the inner ribbon were joined by the magnetizing spiral, c, containing a needle, and a discharge from a half gallon jar sent through the outer ribbon, the needle was strongly magnetized in such a manner as to indicate an induced current through the inner ribbon in the same direction as that of the current of the jar. This experiment was repeated many times, and always with the same result.

800. "When the ends of one of the ribbons were placed very nearly in contact, a small spark was perceived at the opening, the moment the dis-

charge took place through the other ribbon.

801. "When the ends of the same ribbon were separated to a considerable distance, a larger spark than the last could be drawn from each end by presenting a ball, or the knuckle.

802. "Also, if the ends of the outer ribbon were united, so as to form a perfect metallic circuit, a spark could be drawn from any point of the same,

when a discharge was sent through the inner ribbon."

803. An analogous inductive influence to that observed in the case of a series of coils, of which the first in the series is made the channel of a galvanic discharge, has been noticed by Henry in the case of wires placed nearly parallel to each other, of which one is made the medium of a discharge from a battery of coated jars. Having, with my assistance, caused two such wires to pass round my lecture room, as near as possible to each other, without touching, the ends of one of the wires being severally inserted in cups of mercury, while the other wire was the medium of a discharge from my battery, shocks were received in the fingers when they were plunged in the mercury in the cups, at the moment of the discharge.

804. It is to discharges of this kind, as I conceive, that the souring of

milk and beer, during thunder storms, may be ascribed.

805. It is not possible to do full justice to the speculations, experiments, and observations of Henry, in an elementary treatise: I therefore forbear to extend this account of them further.

806. As respects the hypothetical explanation, I have resorted, in common with Henry, to the conventional idea of electrical currents inducing other currents; but it will be hereafter suggested, in treating of the theory of electricity, that the phenomena may be the effect of waves of polarization

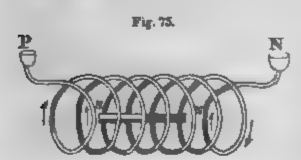
which, in a state of excessive condensation, is conceived to come to space intervening between ponderable metallic particles, and at the metallic in a state of greater or less density to pervade the creation.

### INDUCTION OF MAGNETISM BY CURRENTS.

807. Soon after Ampère, in observing the reaction in tween electrical currents, made the first step in the base of science last treated of, discovery was made of the power of such currents to impart magnetic polarity to a cut if wire, and of rendering a needle situated transversely to the current, magnetic.

808. This power of imparting polarity, was produced increased, when the wire wound into a helix, was made a include the needle. By these means the influence of the current was reiterated, as in the instance of Schweiger's gal-

vanometer.



represents a right spiral of copper wire, containing and of soft iron. The ends of the wire terminate within small cups of mercury. Thus truated, a discharge of the gal-

vanic fluid through the spiral cannot be made so transcutly as not to impart magnetism to the bar. The current being from the cup, P, to the cup, N, the poles of the not will have the character indicated by the initials, S and N: but either reversing the current, or substituting a left spral, would reverse the character of the magnetism communicated to the rod.

810. The polarity given to the needle, when the current flowed one way, was found to be the opposite of that imparted when it flowed in the opposite direction. The current being conceived to proceed in the wire from positive to negative, the end of the needle pointing the same way, as the flowing of the current must receive north polarity, the other receiving consequently south polarity; but reversing the direction of the current, the polarity must be reversed. The polarity varied also, as the helix was wound to the right or the left, so that a left helix being used at mentioned above, the currents would have effects the opposite of those produced by a right helix.

811. In fact the same wire was found to acquire differ-

ent polarities, when different parts of it were made to occupy a succession of cavities in different helices, some wound to the right, the others to the left. A needle was magnetized when situated on the outside of a galvanized helix, but -much less powerfully than when included discharges of "statical electricity were found competent to produce magenetism, when substituted in the same processes, for those . of galvanism, but the power imparted was comparatively : **very** feeble.

812. Mr. Ampère, besides other interesting discoveries which my limits will not allow me to illustrate, and which it would be impossible to describe satisfactorily, most ingeniously caused the ends of a wire forming a helix to return through the axis of the helix, without touching it, to the middle, and, passing them out of the helix, one on one side, the other on the other side, they were made to form an axis, on which the whole might revolve. The different ends of the wire, thus arranged, being made duly to communicate with a galvanic battery, the helix was influenced **by** a magnet, as one magnet is affected by another.



813. Two coils wound one to the right, the other to the left, as in figure 76, and made the medium of a galvanic current, in the usual way, will impart opposite polarities to two bars or rods of iron simultaneously introduced, one of them within the coil on the right, the other within that on the left.

polarity of the outer extremities will be similar, and the same is true of the ends near the centre. This may be shown by withdrawing and presenting them to the poles

of an ordinary magnetic needle.

814. In this way if one bar be introduced so as to occupy the cavities of both spirals, it will have three poles, the middle having only one polarity, which will be the opposite of the similar polarity of the extremities.

### Of the Electro-Magnet.

815. A rod of iron, magnetised by a galvanic current, as above illustrated, is called an electro-magnet. Professor Moll perceived, that if a bar magnet was more powerful in the horse-shoe form, the same ought to hold in the case of an electro-magnet. Hence he was led to contra an electro-magnet resembling that represented in the

joined figure.

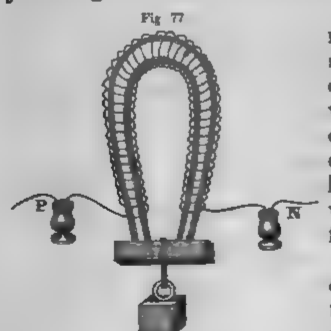
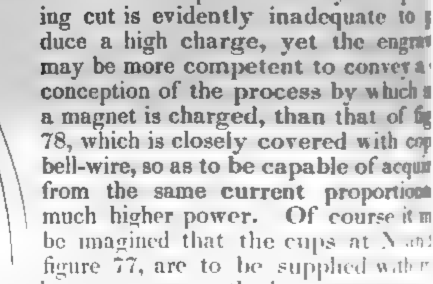


Fig. 78.

816. It consists of anim rod in the shape of a lor shoe, on which copper of the size used for bellet vered by silk, is wound, in one end to the other. electro-magnet thus 🚥 held seventy-five pom while weighing itself five pounds.

817. The quantity of employed in the constr tion of the electro-mag represented by the pres



cury, so as to make a connexion with the extremities a Yet I prefer set screws, for effect galvanic battery. such connexions.

### Of Henry's Electro-magnet.

818. The results obtained by Moll were eclipsed those accomplished by our countryman, Professor Jose Henry, of whom mention has already been made.

819. Of his celebrated electro-magnet, the following a count is extracted from Silliman's Journal, vol. 21, 183

page 400.

820. "A soft iron bar, two inches square, and twen inches long, having the edges rounded, was bent into t form of a horse-shoe. Five hundred and forty feet of co per bell-wire were wound round it, in nine coils of si

sfeet each. These coils were not continued from one end sof the magnet to the other, but each of them was wound round a portion of the horse-shoe about an inch in length, deaving the ends of the wires projecting, and properly numbered. The alternate ends were soldered to a copper cylinder, and the others to a small cylinder of zinc, containing only two-fifths of a square foot, and forming a voltaic arrangement with dilute acid. When the armature of soft iron was placed across the ends of the horse shoe, it was found capable of supporting 650 pounds: an astonishing effect for so small a battery, which required a charge of only half a pint of dilute acid. With a larger battery, the weight sustained was 750 pounds, which seemed to be the maximum of magnetic power that could be developed in that bar by voltaic electricity. It is remarkable, that when the ends of the wires were united so as to form a continuous wire of 540 feet, the weight raised was only 145 pounds."

821. Another magnet was made by Professor Henry,

which held upwards of two thousand pounds.

822. Subsequently magnets have been made adequate to

hold three thousand pounds.

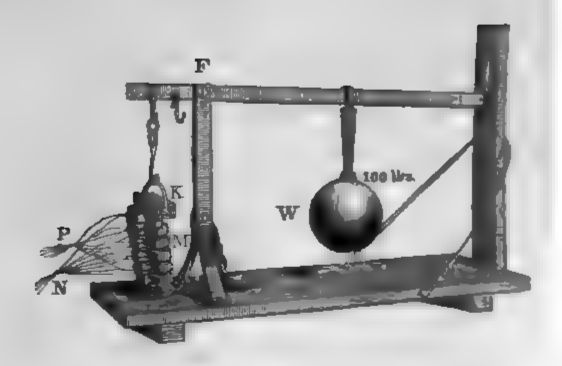
823. As soon as the current through the coils of an electro-magnet is arrested, the keeper, if loaded, falls off, since the magnetic power which remains, is extremely feeble. The rapidity with which the magnetic polarity is produced, destroyed, and reproduced, or reversed by shifting the direction of the current, is very remarkable.

824. The following figure represents a modification of Henry's electromagnet, as constructed by me, soon after the account above quoted was

published.

825. M is the electro-magnet, consisting of an iron cylinder, of 12 inches in diameter, and 24 inches in length, bent so as to form a U. It has six coils of sixty feet of covered copper wire on each leg. K, the keeper, of which the sectional area should not be less than that of the bar of which the electro-magnet is formed. The keeper is suspended from a steelyard, divided into parts, each equal to the distance between the hook supporting the keeper and the pivot of the fulcrum at F. The iron globe W, weighs 100 pounds, and when slid to the fifth division necessarily acts on the keeper with a force equal to five times its weight. When excited by a calorimotor of two pairs, such as has been described, fig. 8, it has been necessary to add additional weight to increase the force to 750 pounds, before the separation of the keeper could be effected.\*

The coils are twelve in number, each comprising sixty feet of copper bell-wire, wound up with thin paper and shell lac varnish, to prevent metallic contact. They



### Of the Rapidity with which Polarity may be Reversed, and the come Rapidity of Rotary Motion thence arising.

826. The attractive power above mentioned as imparted by an electrone to a cylinder or bar of iron in the U form, is rendered not prising by the inconceivable minuteness of the time requisite for the duction, and reversal of the polarity on which it depends.

827. The alternations of attraction and repulsion resulting from the versal of polarity, has been an object of much attention and expension trivance, with the view of obtaining an advantageous moving permachinery.

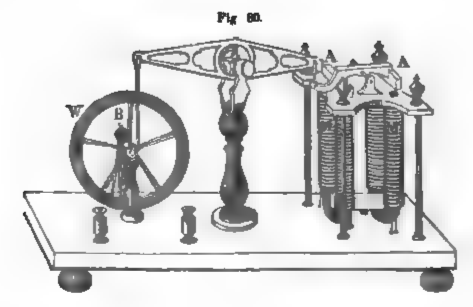
829. Previously to the career of Professor Henry, it was known to most transient duration of a galvanic current passing through a would magnetize, or reverse the magnetism of an iron rod contained the helix. This susceptibility of opposite magnetization was more presented to attention by Henry, and may be illustrated by a latterine, of which a description is subjoined.\*

were severally wound upon an iron cylinder of the same diameter as that electro-magnet, previously surrounded with thick paper so as to increase the little. In consequence of this precaution the bore of the coils was sofficient cious to allow them to be transferred with ease, from the iron cylinder to the of the magnet. Six were thus affixed to each leg. In order that such a set should cooperate, when all are wound one way, so for instance, as to firm not all the beginnings must communicate with one pole, all the endings with the To facilitate this, the analogous extremities are soldered together as represed and N. The same rule must be observed if they be all wound to the est a right and a left coil, may be made to cooperate in producing the same piff each beginning is applied to the same pole as each ending. By a beginnean that extremity at which the coil is begun to be wound by an each course, that extremity at which the winding terminates.

#### \* Reciprocating Engine

Figure 80, may afford a good idea of a reciprocating machine working a; principle suggested by Henry. The polarity of the two upright electro-magn

829. Davenport's Machine, which drew so much attention, added no-The change from an alternate to a rotary thing to our knowledge. motion, was an easy step made independently by Richie and others, as well as himself. As respects its efficacy as a moving power compared with an alternating motion, it was a step towards the rear, which any one well acquainted with the nature of the peculiarity of the powers of electro-magnets and with mechanics, would not have taken. In my opinion, the great defect of this power is, that its force is exerted through a very minute space. For its advantageous exertion it is requisite that the surfaces of the armature and those of the poles of the electro-magnet be parallel and exactly opposite. These objects cannot be well obtained when either the armature or the magnet are undergoing a rotary motion. In that case, when the position of the masses actuated by the opposite polarities, is most favourable for the exercise of their reciprocal attraction or repulsion, their position is least favourable for the production of circular motion. It is in



M, is established or nullified alternately, as the fly-wheel in revolving makes a communication with the coil of one or the other, by a mechanism called the break piecs. The little cylindrical rod which forms the axis of the fly-wheel W, is filed away at two places of about a quarter of an inch in width, on opposite sides of the rod, leaving between them, a part which is not deprived of its rotundity. Of three springs proceeding vertically from beneath the base board, so as to be held firmly by it, one communicates by a wire beneath the board with the set screw on the right and through it with a pole of a battery. Pressing at the same time on the intermediate round portion of the axis, it keeps it constantly in the same electrical state as the pole of the battery with which it communicates. Each of the other two springs communicates through the wire of one of the believes Ma M, with the left set screw, and through thus with the other pole of the battery. The axis, as above mentioned, being in constant communication with one pole, the current alternately passes through the helix of one or the other electro-magnet, accordingly as the spring with which it communicates is, or is not, in contact with the axis in consequence of the alternate presentation at every semi-revolution of the portion which has been removed by the file or that which has been left in statu quo. As the portions in the last mentioned state, are on opposite sides, the contact of the springs with the axis, is alternate. Thus each magnet is alternately charged and alternately free. Consequently two keepers of soft iron associated with each other, and with a beam crank and fly-wheel, by being alternately attracted and liberated, undergo an alternate motion which is imparted to the beam. This motion by means of the crank causes the fly-wheel and its axis to revolve, and thus gives efficacy to the mechanism of the break piece.

It follows that the machine continues in motion as long as the galvanic current is

applied.

The analogy between the mechanism of this apparatus, and that which is resorted to in steam engines, is self-evident.

the vicinity of the " dead point," so called technically, that there a the out

intense reaction.

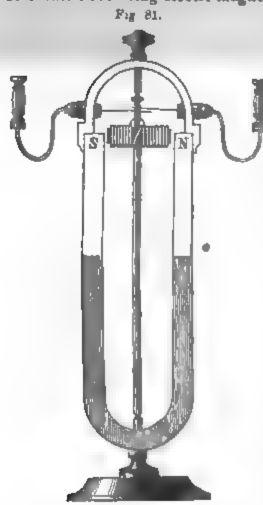
In both, the alternation of polarity by which the beam was set to make agreeably to the apparatus last described, was made to act upon at magnet turning on a vertical axis between the jaws, either of a retained magnet or of an electro-magnet. The former was employed by first the latter by Davenport. I have been under the impression that first it not make due acknowledgment to Henry, for the bunt afforded by alternating beam engine. Yet De la Rive alleges that he are to retain ratus, in 1828, which had been contrived by Richie, to revolve by the ternation of polarity, resulting from changes in the direction of the parameters.

# On the great Advantage of Electro-magnets as the Manu finparting Permanent Magnetism.

831. One of the practical advantages arising from the art of making electro-magnets is, that we are enabled to impart instantaneously the most powerful magnetism to steel bars. The mode of employing for this purpose a electro-magnet, does not differ from that already described

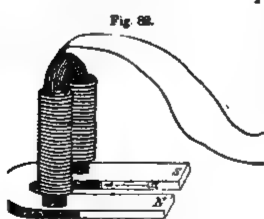
### \* Richie's Revolving Electro-magnet.

The following figure will give an idea of the apparatus which in Europe a task as Ruchie's revolving electro-magnet.



Two cups of mercury, or two set series, are supported severally on such set of the arch of brass, by means of wires transmit each in one of the springs of a poir thate (767) By these means the hear a tro-magnet is subjected to a come. battery, of which the direction is but - " often as the electro-magnet as an in semi-revolution If, while extern ... current the electro-magnet be nade 123 through a quadrant, so as to be at 🛫 😘 🥫 to the position in which it is represented as of its poles will be attracted by corner repelled by the other pole of the land between the legs of which it is place. as soon as I berated, it will seek ' i mag to. But in being thus impered it is rotary force which carries it so far and the current reversed by the pre entry is now impeded to reverse its part. half a turn encounters a change to a from the same causes as at first The area tion ensues and endures so long as teris supplied I have seen the elect. supported horizontally ipin the vertice as represented in the preced no hours and so rapidly as to make a hum - grant to assume the appearance of an obtain spirit or turnip-shaped disk

n using the permanent magnet for the same purpose (729). n either case, however, when the magnet to be charged is of the U form, the induction is more powerful if an electroagnet having the same distance between the legs is emloyed. As the coil may be made loose enough to slip on and off (825), rods of iron in the U shape, may be provided o as to suit the various permanent magnets required.

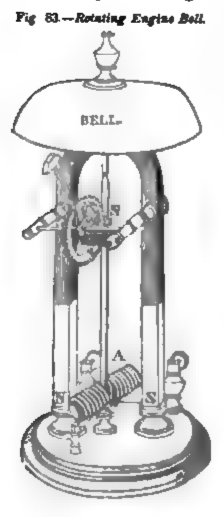


832. Figure 82, will serve to illustrate the proper position of the electro-magnet relatively to a steel U bar to be magnetized, as when first applied. It is to be slid from the position in which it is represented, towards the ends of the steel U bar, two or three times; and finally being

rought into the situation of a keeper, let the keeper be pplied before the magnet is slid off.

### Dynamic Gold Leaf Galvanoscope.

833. The figure of the gold leaf galvanoscope was introduced, in treating



of the galvanic circuit, in order to give the student an idea of the means which it affords of detecting a galvano-electrical current. The explanation of this instrument, requiring a reference to the laws and phenomena which fall under the head of reaction of magnets with electrical currents, was postponed in order to be given here, the figure being of necessity reinserted in the next page.

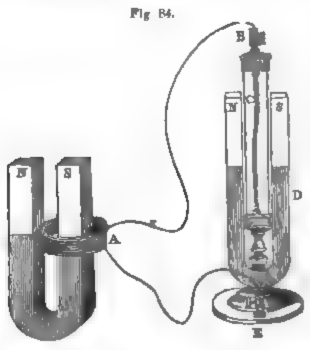
834. Between the legs of a U magnet, fig. 84, a glass tube is placed, to each end

Figure 63, represents the revolving magnet, as associated with a bell, by means of a wheelwork, like that of a clock. An endless screw 8, on the axis acts upon the teeth of a wheel, which as it revolves by means of a projecting pin, lifts the handle of a hammer until it arrives at a due elevation, where it is released from the pin. Then by the reaction of a spiral spring, the hammer is make to strike the bell. As the wheel has sixty-four teeth, it takes sixty-four revolutions of the magnet to cause one stroke. In some instances the number of strokes in a given time being ascertained, the revolutions were found to amount to more than one hundred in a second.

as to be concentric with the axis of the tube. Between the forces to supported and another similar pair offixed to the lower cap of the satisfact of gold leaf is extended, of which one of the two unders body each forceps, as to make it occupy the central space about the unit tube between the forceps. The caps communicate with set screen, which wires are to be affixed, to complete any circuit, which is to the wife ably to the law above stated (626), endeavour to quit the posters which holds, in order to take the tangential attitude, and while their street escape from between the jaws of the magnet, inclines one way or to other, in a plane at right angles to one passing through the area of the jaws (740).

# Of the Induction of Electrical Currents by Magnetin. Magneto-electrical Induction.

835. The influence of electric currents in inducing netism, having been fully demonstrated, the celebrated h raday was led to inquire whether the converse might m be attainable. Accordingly he has ascertained, that ele trical currents may be induced by magnetic action. 4 on the one hand, an electrical current circulating through a coil of wire will produce magnetism in an iron in situated within the coil, so, on the other, if a bar pre viously magnetized be alternately introduced into the col and withdrawn, currents will be induced in the coil at each ingress and egress of the bar. To the currents thus the duced, as well as to those already described as resultafrom the inductive influence of the current in one lesproducing a current in another in its vicinity (781), the appellation of Faradian has been given in honour of the discoverer.



ence of a magnet, in causar an electrical current, may be illustrated by the process which the adjoining figure exemplifies. It will be processed, that the gold leaf of the galvanoscope, alread described (757), is made, a represented in the engravia, to complete the circuit of coil of covered copper wire which has been so wound a to leave about its axis a very series of the control of the circuit of the country of the coun

Ecant space sufficient to receive either of the legs of a U

magnet.

837. In putting the coil into the situation in which it is represented, the gold leaf will incline one way; on removing the coil, it will incline the other way: moreover, the flexure which arises on putting the coil on one leg, will be the opposite of that which will ensue when it may be put about the other leg. Yet removal from one leg, and placing it about the other, will be productive of a similar flexure. While the coil is stationary, the gold leaf is not affected. It is only during the changes of its situation relatively to the magnet, that any electric impulse is given adequate to the production of a transient current (633).

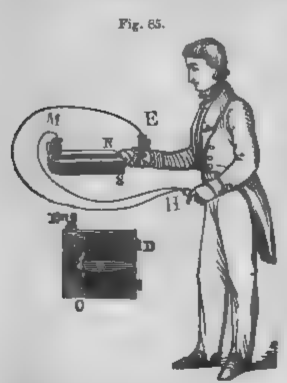
838. It has been pointed out that each end of the keeper of a magnet assumes a magnetic polarity opposite to that of the pole with which it may be in contact, but that the magnetism thus acquired, ceases on the removal of the keeper from the magnet. If we suppose such a keeper to be surrounded by a coil of insulated wire, we have only to remove it from its appropriate position, and restore it alternately, in order to produce a Faradian current within

the coil at each change of position (835).

839. Under these circumstances, the magnetism of the keeper is destroyed as often as it is removed from the poles, and restored as often as it resumes its appropriate position; and an alternate accumulation and subsidence of the magnetic fluid within the keeper, is supposed to produce Faradian currents by its reaction with the electric matter in the wire.

840. The current produced by this last mentioned process was detected by Faraday, by causing the ends of the wire coiled about the keeper to communicate with the terminations of the coil of a multiplier, at the moment of removal from the poles of a powerful magnet. But of the current thus detected, indications were soon after obtained by the production of sparks. This interesting result was first attained by the distinguished Italian electricians, Nobili and Antinori, by causing the ends of the wire coiled about the keeper to break contact with each other through the medium of an amalgamated surface, in consequence of the jar arising from the lifting of the keeper from the poles of a magnet. Professor Forbes, of Edinburgh, produced

the same phenomenon shortly afterwards, by employing the same process with a powerful loadstone.



841. This experiment is dominated by the adjoining figure. The arms is a U ber of soft from, and is second ed by a fine wire coal of great and It may, advantageously, be fifices 😂 dred feet long. Of course a last customary covering of cotton or A section of it, and its case, a ransented at E, O, D. One end of the wat is soldered to the armature and the 🖚 The other end of the wire is sodewill the set screw, E, insulated from the by ivory. H, held by the operator, at metallic handle with two wires. 🖦 🛊 attached to the set screw, E, and the communicates with one end of the the other communicates with the of through the set screw, M, communated with the magnet, and of course with it

keeper is applied to the magnetic poles, it becomes a magnet (733; who separated, it returns to its previous state. The internal change which to keeper undergoes in acquiring or losing polarity, is productive, by the tion, of cotemporaneous currents in the coil. The resiliency of the pile ized matter in the keeper, when it is suddenly put on, or pulled off from the poles of the magnet, acts like that of a galvanic circuit when sadding opened or closed (775). Hence, simultaneously with the removal, a section and a spark is perceived by the operator, who, when the sadding through the magnet is suspended, becomes the only medium through the resilient secondary current (779) can pass. The cure it is the case to the right hand, and through his left, and the ware the read of the coil.

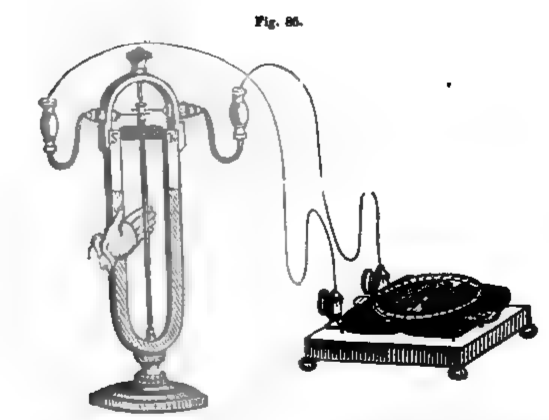
842. The changes of the relative situation of the keeper and magnet which the current within the coled wire is excited, and arrested in the versed alternately, were much facilitated by a mechanism contributed Pixii, of Paris, which caused the rapid rotation of a magnet restrict takeoper, surrounded by a coil of wire. But, by Saxton, an observe the vantageous improvement was made, in causing the keeper to revoke the stead of the magnet.

843. In consequence of the greater rapidity with which the reversitathe positions of the keeper could be accomplished by the aid of banks as multiplying wheels, the magneto-electric machine thus contrived by Parand improved by Saxton, not only produces deflections in the neede. as a succession of sparks which seem to have no intermission, but becausevere shocks, the ignition of wire, and chemical decomposition.

844. If Richie's revolving electro-magnet, as modified by Page \*1', a connected with a galvanometer, as represented in the adjoining figure, a causing the axis and electro-magnet which it supports to rotate, at ever semi-revolution in the needle will be observed a deviation, which will also nately be opposite in direction.

1

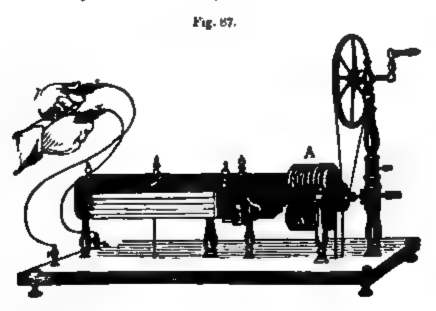
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845. This phenomenon is perfectly analogous to that to which attention is in the next place to be given, in which a current is produced at every alternation of polarity of an iron keeper, consequent to the reversal of situation relatively to a permanent magnet.

Of the Magneto-electric Machine.

**846.** The following figure represents Saxton's magneto-electric machine, well constructed by Daniel Davis, jr., of Boston.



847. A, represents the armature of this apparatus, supported on an axis, so as to be as near as possible to the terminating surfaces of the legs of a compound magnet. Of this axis one end reaches between the legs parallel to and equidistant from them, to a socket in the arched part. The other end of the axis, furnished with a pulley, terminates in another socket beyond the armature; so that, by means of a wheel and band, it may be made to revolve with great celerity.

848. The iron of the armature forms three sides of a rectangle. On the two opposite sides, 2500 feet of copper wire, of a grain to the inch in

weight, are wound so as to involve each side of the rectangular keeps as coil.

849. It has already been explained, that an armature becomes a capt by induction, so long as it is applied to its imagnet; but that as pure ceases when it is removed, or may be induced in the opposite was it is

reversal of its position relatively to the magnet (733).

850. Now these changes all ensue during every semi-rent and during the shifting of the polarities a corresponding change is to the surrounding coil, equivalent to what is called an electron of the The coils, when their ends are in communication, have product at them alternate circuitous discharges, having all the attributes of the start occur within the active galvanic circuit.

revolving electro-magnet, fig. 81. In that apparatus, alternations of constrained polarities are produced by alternations in the direction of constrained current; in the magneto-electric machine, alternations of experience currents are produced by alternations of opposite magneto-posities. This must be evident from the preceding illustration (529).

the axis, is connected with one of the extremities of the wire sure the keeper, the other end of this wire communicates with one of the screws of the board, a spring of metal attached to an upright round brass, plays upon the teeth of the revolving toothed wheel, and cases cates with the other set screw. Any portion of the naimal frame, and made to complete the circuit between the set screws, will receive a declarising from a resilient current, at each interruption of the circuit cased the intermission of contact between the spring and teeth, as a keep in one tooth to the other.

853. When cylinders of copper, soldered to the ends of the crist rounding the keeper of a powerful magneto-electrical machine, are graph firmly with the hands previously moistened, the most powerful makes able during the action of the machine to relax his hold, he is previously mercy of the operator controlling the wheel of the machine.

854. Experience has shown that, for the production of the start wire coiled about the keeper should be of comparatively great and fifteen hundred feet, but for producing limition, should be much start and not more than eighty feet in length. The propriety of not have said ence of length, Mr. Saxton assures me, was first suggested by training

applies the keeper to the magnet on one side, near the ends of the state that in his electro-magnetic machine the magnet is vertical, where a su ton's it is horizontal. A priori, this would seem to promise no always and yet machines thus constructed have proved at least as power'd as it others. The only advantage which I can conceive of, as resulting in a direction recall to confirm the polarity of the steel magnet.

856. The magneto-electric machine has been resorted to as a remove of nervous, rheumatic, and paralytic affections; and at all events may be use as a medicina mentis, which probably will do no harm, even what does no good. It has a great advantage over common electrical machine in being independent of the weather, and is preferable to voltaic apparatus in not requiring the use of corrosive liquids. The apparatus may be as ried in a box, without detriment, by land, as well as by water.

### Magneto-electric Electrolysis.

857. The process by which the decomposition or electrolyzation of water, is accomplished by the currents induced by a magneto-electric machine, may be understood from the annexed engraving. Two wires proceeding from their solderings to the set screws A and B, may be observed to reach vertically into the trumpet-shaped orifice of the tube G, which is situated over them, and which must be at the outset replete with water.

858. It has been mentioned that to obtain the intensity requisite to severe shocks, the circuit is abruptly broken by a toothed wheel, while otherwise it would continue until the keeper should be off of the magnet. But this secondary current is not sufficiently enduring to be competent for electrolysis. In this respect it resembles dis-

narges of statical electricity. Moreover, the current sould not alternate in direction, as this would render it apossible to obtain the different elements separately.

859. In order to cause the electricity to flow always the same direction, a pole changer, like that already escribed, must be employed. For the same reason at a current otherwise uniform in its course, may by is pole changer be made to flow alternately in opposite ays, so where a current is alternately opposite, the pole nanger may render it uniform in its course. The change position of the silver segments which convey the curnts to the set screws, to which the electrodes of the batry are affixed, and the change in the direction of the dislarges taking place simultaneously at every semi-revoluon, if a segment begin with the reception of a positive scharge, it must receive the same at the end of a half rn, and of course the set screw with which it communiites and any conductors thereto attached. qually true of the segment which receives at the outset a gative discharge.

860. To effect the collection of the elements of water parately, it is necessary to employ two tubes O, H, one of

In this case as in others, the ratio of the bulk of the hy gen to the oxygen, will be indicated by the space retively occupied by them. With a good machine, to each of a mixture of both gases, may require tween five and ten minutes.

tric machine, various saline solutions may be composed. For this purpose a tube in the confidence of the letter U may be employed, figure to pledget of cotton cloth may be made to oct the bend, as a partition. The solution occuped by appropriate tests. Any salt with a metallic number which does not decompose water per se, will depose metal on the negative electrode. Hence this process be employed in gilding or silvering by electrolysis.

### Improved Magneto-electric Machine.

662. Since the preceding descriptions were written, I have receive Mr. Davis, a magneto-electric machine, upon a construction which dently new, and of which the effects are very energetic.

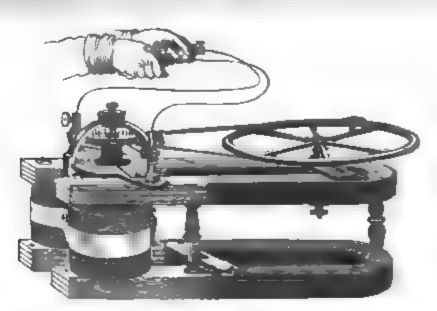


Fig. 90,

863. It will be perceived, that in Saxton's machine, \*0, the keeps soft iron bar so bent as to form three sides of a rectangle A. B. C. of the two opposite sides B, C, may be designated as legs. Each leg is enciral helix of covered wire. The end of each leg is acted upon by one parameters of compound magnet, the side intermediate between the legs serves of be a mean of communication between them, having no ability to be the magneto-inductive influence.

864. But in the new instrument, a second compound magnet is as

Example the place of the inactive part of the rectangular keeper; so that two important in possible polarity by their application, at four extremities of two bars, instead of two extremi-

ies of one, equivalent to two, usually employed.

1 865. Of course each of the iron cylinders which takes the place of the least of a rectangle in Saxton's construction, are presented to poles endowed with an opposite magnetic power. The north pole of the lower magnet is thus made to co-operate with the south pole of the upper one, and in like manner the south pole of the upper magnet with the north pole of the lower magnet, just as is the case when a keeper is in its appropriate place, each of its ends being in contact with one of the poles of the same magnet, the influence of these has a concurrent effect in producing an opposite polarity at the extremities of the keeper.

stance of a statement in a letter from my sagacious and much esteemed friend and pupil, Dr. W. F. Channing. The unmitigated shocks from this machine are insupportable. When the wires which break the shocks are removed, the current becomes sufficiently uniform to be competent for electrolysis or imparting magnetism to iron included in a long helix of fine wire comprised in the circuit of the helices of the machine. When sent through a circuit of a mile, the current from this machine, was found abundantly

competent to work the telegraph of Professor Morse.

### Of Ferro-dynamic Induction.

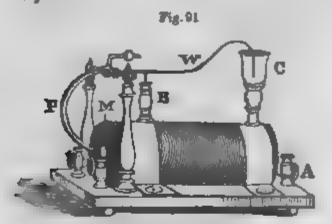
867. I venture here to use an epithet for which I have no authority, so far as respects the employment of the Latin word for iron to designate the property of a mass of this metal to heighten the inductive influence of a coil by which it is magnetized, whenever the primary current is set in motion, and de-magnetized when this current is arrested.

## Of Callan's Faradian Apparatus, and Page's Callan and Faradian Electrotome.

sor Callan constructed a very powerful apparatus, by surrounding an iron cylinder with two coils of covered wire, one within the other, the inner coil being coarse and short, the outer one long and slender. When the inner coil was made to form the circuit of a voltaic battery, the other was found to give severe shocks or brilliant sparks, at the moment of interrupting the circuit.

Raradian principle. One of these is extremely interesting and ingenious, as it is so constructed as that the magnetic attraction of the included iron, arising from the influence of the circuit, attracts a lever from the position in which it forms a necessary portion of the circuit. The circuit is thus broken, but as the lever falls into its previous position, the circuit is renewed. Each time that the removal of the lever takes place, a shock is received by a person duly communicating with the terminations of the finer wire; and

a brilliant spark appears as often as the lever quits the surface of the



10. The apparatus a to, is represented by the ing figure 91. A code of covered copper wire about a bundle of strught iron wires like kning a The heirs thus coests surrounded by a bein covered copper wire of a cone thousand feet, of whends are soldered to exp

screws inserted into the base board on the right. These set screws attaching severally one end of each of two wires, having a hadic free end. One of the ends of the inner coil is attached to the aut at on the right, while the other end of this coil is soldered to a bras which secures the coil and supports the brase cup B, holding m The cup C similarly occupied, is connected with another set screw ponding to A, in size and situation. The wire W, which plays on zontal axis by means of a spur and its right hand termination, exalt conducting communication between the cups of mercury B and C, t is kept down by its own unresisted weight. It carries, however, a f iron at the left extremity, within half an inch of the bundle of course As soon as the set screws on the right, which communicate with the coil, are made to receive the electrodes of a galvanic battery, the ba wire acquires an intense magnetic power, and attracting the iron a to the wire beam W, lifts the right end out of the mercury in the This breaks the circuit, to which the magnetic attraction of the coland the weight of the wire beam no longer counteracted, cames & back to its primitive situation. The magnetism is thus renewed, a circuit is again interrupted, by the lifting of the end of the ware harthe surfaces of the mercury in C. These alternations are accor. with surprising rapidity, a brilliant spark being given as often as " tact with the mercury is broken. Meanwhile a person grasping to dles attached to the ends of the smaller coil, will receive a shock at breach of the circuit, through the coarser cod. The shocks this rienced with an apparatus of a moderate strength and size, white very severe.

971. Of the shocks resulting from the inductive power of one cupon another, an account has been given. In that case, it seemed as effect of a larger coil upon a smaller one, is to cause a degree of a polarity commensurate with the weight and length of the inductor but that in the discharge of this polarity, the intensity was inversely

sectional area of the wire forming the coil.

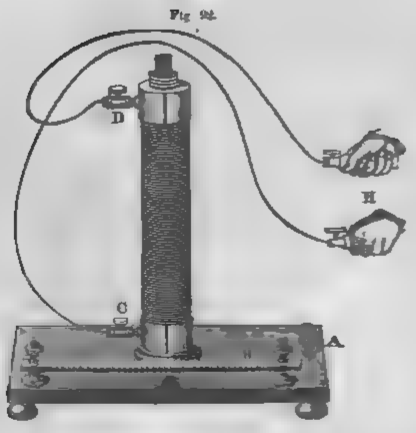
872. But in the secondary current produced in Callan's agrama Page's modification of it, which I have been describing, we have at the reaction of all the electro-polarity of the coil, an additional first that of the magnetic polarity of the iron. The tron in the electrical dergoes the same magnetic change as that in the armature of the magnetic machine, although from a different cause. In the one case duces a current, in the other it merely promotes a current, which exist independently of its aid.

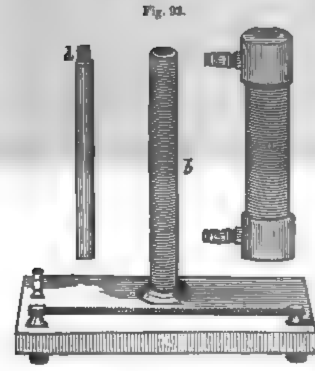
Decomposition aided by the Influence of the Resiliating Current of an Electrotome.

873. M. De la Rive has devised an apparatus which he terms "Condensature Voltaique." It comprises a small bar of iron, on which is closely wound thick copper wire, covered with silk. Through this wire the current of a single voltaic pair is made to circulate, while by means of a little electro-inagnetic apparatus acting like Page's electrotome above described, the circuit of the current is broken and reversed. An apparatus for the decomposition of water being included in the circuit, it is found that the electrolytic effect of the voltaic pair employed is nearly doubled; while a bright light is afforded by interposed charcoal points. The voltaic condensor succeeds well only with a battery on which two electrolytes are used, as in those of Daniell, or Grove. For its success it is necessary that two chemical actions be going on at once within the voltaic pair employed with it. The wire wound around the bar should also be large in diameter, and of moderate length. The action of the apparatus is referred, by its inventor, to the depolarization of the platinum electrodes in the decomposing cell. Since its introduction into the circuit, however, produces precisely the same effect as would result from an addition to the number of pairs, a part at least of its efficacy must be referred to the efficiency of the iron as a reservoir of the polarity of which the resiliating discharge (775) takes place, with a velocity greater than that with which it is accumulated.

# Page's Apparatus for Regulating the Energy of the Resiliating Shocks of a Magneto-Electrotomic Apparatus.

874. The great power of iron in increasing the capacity of a coil, upon Callan's construction, to give shocks, is beautifully illustrated by an apparatus made by Davies, under the direction of Doctor Page. Agreeably to the construction of this apparatus, two different coils, one of fine, the other of coarse wire of Callan's apparatus are separable. Figure 92, represents the instrument in question, as when the parts are duly associated. In figure 93, it appears as when the coils have been taken apart. The external helix, a, is formed of fine covered copper wire, not being less than one thousand feet in length. The ends of this coil are soldered severally to the set screws C, D. The inner coil consists of not less than three strands of coarse copper wire. The ends of these strands, which terminate in the same direction, are soldered all to the set screw A, on the base board. The other ends of the three strands are soldered to the serrated





steel bar B: when these cods situated one within the other, 📟 🗈 figure 92, on subjecting the min coils to a galvanic current by taching one of the electrodes wa set screw, the other electrode build in contact with B, on drawing end of the wire, forming the trode, over the teeth of the serrain bar, a brilliant spark w lob 🗝 🖘 the wire leaps from one was " the other. If, during it a prome wires with metallic hapters # 2 tached severally at C and D, 130 son holding the handers w. --for every spark a perceptable story The addition of a single war or knitting needle will increase thest < sensibly, and by the further accu-

of such wires the shock will be greatly increased, so that if the galvin battery employed has only a square foot of zinc surface, the sensation of comes such as few are willing to bear. A rod of iron is not quite so that cious as the bundle of wires of equal size. The inductive influence of the wires is not intercepted by their inclusion in a glass tube, but is much in paired by inclusion in a brass tube, in consequence of its diffusive power lowering the intensity. This is consistent with the experiments of Hear with flat coils (799). The brass tube, like the metallic sheets, interposes closed and perfectly conducting circuit.\*

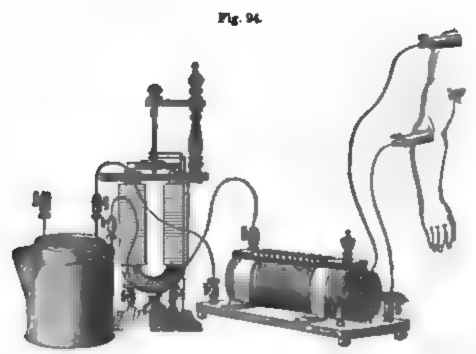
A more convenient form for the application of these shocks as a remedial age is displayed by figure 115, by which the mode of operating is fully illustrated. Their circuit with the battery is established by means of the wire reaching from the s

### Of Thermo-Electricity.

875. When the needle of a galvanometer is deflected by the proximity of any body or set of bodies, of which different portions are of different temperatures, the phenomenon is ascribed to a thermo-electric current in contradistinction to those produced by galvanic apparatus. These have been distinguished by some electricians as hydro-electric currents, as in the necessity of water to their existence, they differ most pointedly from the currents above mentioned, which owe their activity to dry heat.

876. A thermo-electric current resembles a galvanic current of great quantity, but as to intensity extremely feeble. Other things being equal, a thermo-electric current flows from the hotter to the cooler portion of the mass. For the advantageous exhibition of this property, the bodies employed should have a great length in proportion to their other dimensions, as in wires or prisms. the prisms of two heterogeneous metals have a conducting communication with each other between an extremity of each, while the other ends are in contact, or united by a slight soldering, on heating the ends associated as last

screw on the base board to one on the battery, while from the other on the battery, a wire proceeds through the coil of an electro-magnet to a set screw connected with the coarse inner coil of the electrotome.



The interruptions of the circuit are effected by an armature, which, by means of Page's pole changer (766), in revolving reverses at every half turn the direction of the current and the polarity of the electro-magnet, which causes its motion. Any part of the human body, the arm for instance, as represented in the figure, when completing the circuit of the outer coil, will be subjected to resiliating shocks. Although, as these occur at every semi-revolution, the shocks are more numerous than when the serrated steel bur is employed, they are less severe.

mentioned, a current ensues from one metal to the other, through the juncture. The current flows always the many, when the same metals are employed. The metal from which the current flows is said to be positive to the other. It should be called thermo-positive, and those which are

relatively negative thermo-negative.

877. The reaction in this respect is of course consum with that established in respect to metals employed a analogous galvanic processes. As the current in a pivanic triad is stronger in proportion as there is greats discordancy in the relation of the metals employed, to a thermometric apparatus there seems to be a series, in which each metal is positive as respects all that follow, and gative, as respects all that precede it. Heretofore beauti was deemed the most thermo-positive, antimony the not thermo-negative of the metals, and hence, until lately, most efficient thermo-electric batteries were made of muth and antimony. Recently, however, German silve has been found nearly as thermo-positive as bismuth, and is preferable from not being so exceedingly fusible. The fusibility limited the deflection of a needle, subjected to t current produced with a pile of bismuth and antimony, 82°; while German silver with antimony caused a delice tion of 88°. German silver, however, while producing with antimony a deflection of 88°, produced under like circumstances a deflection of 85° with silver, brass, iron, page dium, copper or cadmium; being positive to all of the With zinc it produced a deviation of  $84^{\circ}$ .

878. It has been shown by Nobili, that cylinders of chip partially baked, are capable of yielding thermo-electric

currents.

879. As in the case of a galvanic circuit, a diversity of the surfaces of the same metal or in the liquids to what it may be exposed, will enable it to act as two metals and to give a current from one to the other; even partially twisting a wire, will cause the twisted portion to act at the negative metal, heat being applied at a certain destance from the twist.

880. In all cases where thermo-electric currents are concerned, no less than where those of galvanism prevail, the circuit must be completed in order to have a current.

881. It does not appear that there is any connexion between the thermo-electric, and galvano-electric relation

metals, and those which exist between them as respects weight, specific gravity or conduction either of heat, or electricity. As heretofore combinations of antimony and bismuth were found most efficacious, the one being pre-eminently positive, the other pre-eminently negative, the idea arose that these metals might be indebted for this superiority to their eminently crystalline structure; but it has been found that German silver, with antimony, is more powerfully positive than bismuth is with that metal, although German silver is a malleable metal, of which the structure is not remarkably crystalline.

882. For the following table we are indebted to Cumming, one of the most distinguished among the cultivators

of this branch of science.

| Thermo-electric | Hydro-electric | Series of conductors, |          |
|-----------------|----------------|-----------------------|----------|
| series.         | sories.        | of Electricity.       | of Heat. |
| Galena          | Potassium      | Silver                | Silver   |
| Bismuth         | Barium         | Copper                | Gold     |
| German silver   | Zinc           | Lead                  | Tin      |
| Mercury ?       | Cadmium        | Gold )                | Copper   |
| Nickel \$       | Tin            | Brass >               | Platinum |
| Platinum        | Iron           | Zinc                  | Iron     |
| Palladium       | Bismuth        | Tin                   | Lead     |
| Cobalt ?        | Antimony       | Platinum              |          |
| Manganese }     | Lead           | Palladium             |          |
| Tin             | Copper         | Iron                  |          |
| Lead            | Silver         |                       |          |
| Brass           | Palladium      |                       |          |
| Rhodium         | Tellurium      |                       |          |
| Gold            | Gold           |                       |          |
| Copper          | Charcoal       |                       |          |
| Silver          | Platinum       |                       |          |
| Zinc            | Iridium        |                       |          |
| Cadmium         | Rhodium        |                       |          |
| Charcoal ?      |                |                       |          |
| Plumbago 🕻      |                |                       |          |
| Antimony        |                |                       |          |
| 000 TC 4 -      | 1-4'           | 1.1.1.1               |          |

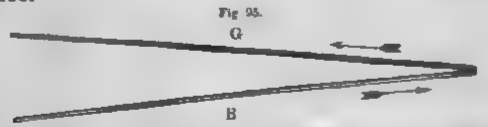
883. If to a platina wire which has been twisted about the middle, a flame be applied at some distance on one side of the twist, the ends of the wire being previously made to communicate through a galvanometer, the needle will indicate that a current flows from the heated part towards the twist. This is ascribed to the twisted part being cooler than the untwisted portion of the wire equidistant from the flame in consequence of a diminution of conducting power caused by the derangement of the metallic fibres during the torsion.

R

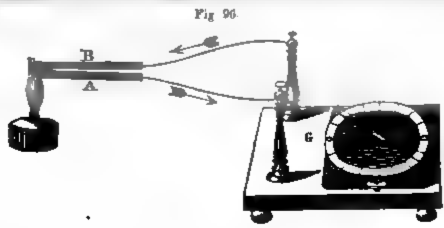
884. The needle will in like manner be affected, if a visionilarly communicating with a galvanometrical cod, is divided in the middle, then each end twisted, and med the twisted ends, after being made red hot, brought and contact with the other. Similar results though lecte, may be obtained by means of copper and solver was That these effects do not depend on chemical reaction who the atmospheric oxygen, has been ascertained by the succeeding under pure oil.

885. If one-half of a ring or rectangle, consisting enter of bismuth or antimony, be heated, while the other half is kept cool by ice, a current will be excited sufficient to affect a needle, although no coil be employed to heighten in

influence.



986. Let one end of a wire of German silver G, be brazed to cot all of a brass wire B, on heating the juncture and making a communicate between their other two ends by a third wire, a magnetic need's duly applied will be affected, as if a galvano-electric current of ferble intensity was circulating in the circuit thus formed. The deviation of the needs is as to indicate that the thermo-electric current thus induced, flows from the wire of German silver to the brass wire.

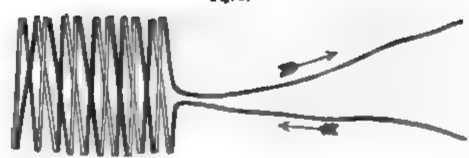


a bar of antimony, exactly alike as to dimensions. Each is represented as a communicating, at one end, with one of the terminations of the wire of a galvanometer. Their other ends are, while touching, exposed to the flame of a lamp. Under these circumstances the needle of the galvanometer will be deflected. Of course the arrows indicate the current to take place, from the thermo-positive metal, to that which is comparatively thermo-negative.

888. Figure 97, represents a series of eleven pairs of German sive and brass wire, arranged in two rows, and soldered as above desembed 4 voltaic order. It is sufficient to heat or cool the junctures on either select produce a current competent to deflect the needle of a galvanoscope, we







h it is made to communicate as represented by figure 96. Even the of the hand is sufficient to cause the needle exposed to the consequent ont to deviate to a perceptible extent.

## struction of a Thermo-electric Pile of Bismuth and Antimony, illustrated by an Engraving.

 Let any number of bars of bismuth and antimony, exactly similar I their dimensions, be buildled together, paper being interposed between lateral surfaces, so as to avoid metallic contact. Let the neighbouring of the heterogeneous bars thus associated be alternately soldered, and nately left free, so that any bar will have only one soldering at each and not be soldered at more than one end to any one bar. hich the bars must be united is exemplified in figure 97. The bars connected, will form one continuous metallic conductor, consisting nately of bismuth and antimony, so that on taking the whole into a galc circuit, all lateral discharge being prevented by the paper, the current ring by a bar of bismuth, would proceed through a bar of antimony, through a bar of bismuth, and so on coming out finally through a bar The metals are in fact alternated like those of the couronne 'asses, excepting that the fluid, instead of being alternately conducted by il, alternately transmitted convectively through an interposed liquid, is eyed from bar to bar, wholly through the alternate metallic junctures. 30. The metals thus associated, exercise, when their alternate solderings at different temperatures, an electro-motive power due altogether to contact; so that the contact theory of Volta, however incompetent to unt for the powers of his pile, seems to be somewhat applicable in the case of the thermo-electric series. It is not pretended, that in the case he apparatus last mentioned, any chemical combination takes place, ss the successive passage of heat from particle to particle be considered 'arranting such an idea.

Fig. 98.

891. By figure 98, a thermo-electric battery is represented, consisting of sixty-three pairs of bars of bismuth and antimony, each three inches in length, three-fourths of an inch in breadth, and one-fourth of an inch in thickness, of course the bars are soldered to each other alternately, as above described, so as to be arranged in a series in voltaic order. They are insulated and secured by plaster of Paris,

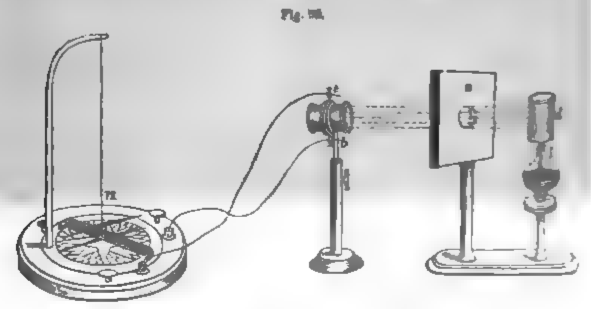
copper case, B, having above them, a space for receiving ice, and below her so situated as to be exposed to an appropriate cast iron plate, presly heated to redness.

92. Two set screws for attaching wires are represented externally, each thich is made to communicate with one pole of the battery. The energy he current is in proportion to the difference of temperature, created in

those solderings of the bars, which are in the different surfaces of the bard which they form collectively, by the application of heat at use exist and cold at the other surface. Of course the greatest power result in applying ice or a cold mixture to the solderings at one of the sures, at at the other as much heat as bismuth will bear from a bor two which is sion. A series thus constructed, caused the electro-magnet of Robert paratus to revolve rapidly. A serrated bar is attached to the front it case, and is connected with one of the poles of the pile. When a ward which one end is affixed to a set screw soldered to the other pole, but the free end drawn over the teeth of the serrated bar, sparks were producted.

### Of Melloni's Thermo-multiplier.

893. By means of a thermo-electric series analogous to that illustrate by the preceding figure, associated with an extremely sensitive galaxies ter, Melloni contrived a thermoscopic apparatus, more capable of decrease minute differences of temperature than any before known. It provides means of making those interesting discoveries, of the difference in the promotion of bodies to the rays of heat and light, to which reference in made under the head of caloric (C, 309). It has been used to assume the temperature of insects and various parts of the animal system.



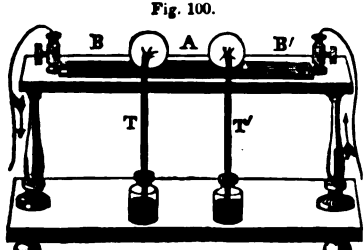
894. The series employed by Melloni, consisted of fifty pairs, about a inch and a fifth in length, and the whole forming a bundle of lattle over a inch in diameter. It was included in a cylindrical brass case, between ward and the bars, no metallic contact was permitted. Wires proceed up ince the extremities of the series were made to communicate with set series but insulated from the brass case by ivory.

895. The mode in which this thermoscopic apparatus was employed my be understood from the preceding figure. The heat of the copper cy at d, received from the lamp l, reaches one side of the thermo-electric sum through a hole in the screen s. The other not being exposed to the influent of any hot body. Under these circumstances a thermo-electric current r produced through the coil of the galvanometer, which by the attachment of its ends to the set screws b, t, is made to complete the circuit between the poles of the thermo-electric series.

896. It is inferred that the difference of temperature, created by the radian heat reaching the surface of the series exposed thereto, will be as the quantit of electricity, and that as the deflections of the galvanometer measure the

electricity, they must also be an index of the degree of heat applied at the appropriate surface of the thermo-electric series. Melloni alleges, that this correspondence between the heat applied, and the deflections, has been ascertained to exist.\*

The Converse of the Thermo-electric Process.



897. The object of figure 100, is to represent a thermo-electric apparatus, in which the phenomena are the inverse of those produced by the thermo-electric batteries, described in the preceding pages. In those a current of electricity is the consequence of a diversity of temperature; in the apparatus in question a diversity of temperature is the consequence of an electric current. The let-

ter A, indicates a bar of antimony so situated between two bars of bismuth B, B, as to have its right end soldered to one, the left end to the other of these bars. Thus the three bars, form one metallic prism, uniform as to shape, though heterogeneous as to the materials, which rests horizontally upon a frame as represented in the figure. At each soldering a little concavity is made sufficient to hold a small quantity of water, and to receive the bulbs of the two air thermometers F and T. The water serves to facilitate the conduction of heat from the metal to the glass bulbs. The prism formed as described, of bismuth and antimony, being made the channel of a galvanic current, in the direction indicated by the arrows, from B through A to B, at the soldering of A to B, cold is produced and made evident by the thermometer T. Meanwhile on inspecting the other thermometer, it will be seen that heat is produced at the other soldering. By reversing the direction of the current, the effect on the thermometer will be reversed.

898. The extent of the elevation above the previous temperature is greater than the depression. This is ascribed to a small degree of heat, which the electric current causes, and which of course adds to the heat on one side, and lessens the cold on the other.

## On Terrestrial Magnetism.

899. The thermo-electric branch of science has a preeminent importance from the light which it has thrown

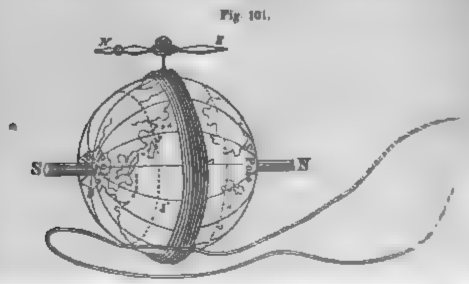
The following are the dimensions of the galvanometer or thermo-multiplier of Melloni and Nobili, associated with the thermoscopic apparatus as stated by Bequerel. The wire is of copper doubly wound with silk, and of about three-hundredths of an inch in diameter. It makes 150 circumvolutions around a frame of about two inches square, and one-fourth of an inch in height. The needles are a little short of two inches in length.

It will be perceived that there are two needles associated so as to form what is called an astatic needle. They are suspended by a single cocoon filament. In the apparatus as described by Bequerel, the cocoon is attached at the upper extremity to a swivel, as in the torsion electrometer of Coulomb, (E, 136,) and the frame may be made to rotate by means of a toothed wheel and pinion so as to adjust the needle to zero, of the graduated arch of ninety degrees, by which the deflections are measured. There is also a contrivance for lowering the needles so as to rest on the frame on which the wire is wound. When thus situated if it be desired to transport the instrument from one place to another, the needles are bound fast to the surface of the frame by a silk ribbon. The apparatus is included in a bell glass to protect it from being agitated by blasts of air, or soiled by dust.

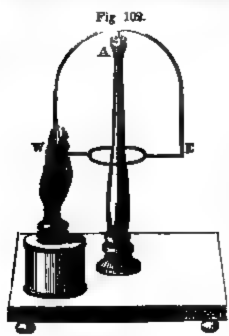
on the nature and causes of the all-important polarity of the earth. This is now ascribed to the successive inteence of the solar rays upon the temperature of the cutt, causing a perpetual thermo-electric current from east to west.

900. It is by the reaction of its poles with this tenetrial thermo-electric current, which flows parallel to the equator, that the compass needle is made to arrange to at right angles to that current, and thus to assume a derection nearly coincident with that of the meridian.

901. In the following figure, a coil of covered copper were affected means of causing an electrical current to circulate about a globa square



senting the earth. A magnetic needle is represented in the posts which it would arrange itself by reaction with the current in the case.

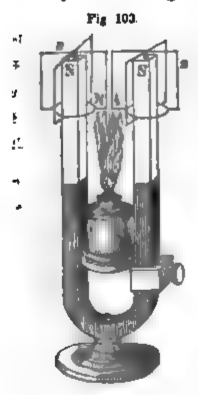


902. The apparatus represented by "..." 101, serves to illustrate the reaction, with magnetism of the earth, of a therm- ber current, produced within an arch of brass w " soldered at its two ends to a ring of German silver. The arch is poised upon a parot at a top of the pillar at A, which the ring enemis The lamp, W, being situated as in the fig = the apparatus must be so placed, that a = passing through the centre of the flame to the of the pillar, will be directed from west these Then, when the flame is applied to the time ture of the arched wire with the ring, a cur rent will flow from the juncture, and of course from west to east through the arch. As we current will conflict with that of the cart. will cause the wire to arrange itself, so as tha

this discordance may be avoided. But a semi-revolution bring ng the other juncture of the arch and ring into the flame, a current through the arch from west to east is again induced, producing another semi-revolution. O course this process will be repeated as long as the flame is applied.

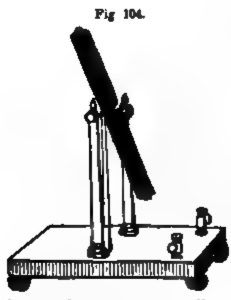


908. Another illustration of the efficacy of thermo-electric currents, is afforded by their displaying an analogous reaction with a permanent steel magnet to that which has been shown to exist between such a magnet and the currents created by galvanism, in the experiment designated as that of ≅ Ampère's rotating battery (757).



904. The frames of wire in the rectangular form, supported on the jaws of the permanent steel U magnet, as represented in the adjoining figure, may consist of silver in rings and platinum rectangles, or the rectangles may be of silver, and the rings of German When the junctures of these frames are subjected to a flame as represented in the engraving, electric currents are made to flow in each from the heated juncture, through the silver wire, to the other junc-The currents, thus incited, conflicting with the polar influence of the magnet, endeavour to arrange themselves so as to escape from conflict. But the only motion which the suspension will permit, is that of rotation, which, by bringing a juncture into the flame at every quarter revolution, causes the effort for a change of position to be repeated as often. Hence the frames revolve as long as the flame is applied.

### Electro-magnets revolving under the Influence of Terrestrial Thermoelectric Currents.



has to the compass needle. represented by figure 104.

905. It has been shown, that an electromagnet, situated on an axis, and subjected to the polar influence of another magnet, will strive to arrange itself so as to bring opposite poles into proximity; but that if the current to which it is indebted for its magnetism be, from the construction, liable to be reversed by a pole changer at every semi-revolution, a rotary motion will ensue. Two instruments have been constructed, in which terrestrial polarity is made to take the place of an artificial magnet, or an electrical current. One of these revolves in a horizontal plane, the other in a vertical plane; the latter having the same relation to the dipping needle that the former The instrument which revolves vertically, is

### Of the Electro-magnetic Telegraph.

906. Though the attempts which have been made to employ electro-magnetism, as a moving power, have hitherto failed, yet another important application of this agent has been made within a few years, in our own country, by Prof. Morse, and in England by Messrs. Barrit and O and Prof. Wheatstone. With the telegraph of Me which has been the medium of communication between the cities of Baltimore and Washington, the public become familiar. The apparatus thus employed, a superior in simplicity to any contrivance elsewhere properation, of which any knowledge has reached the or try. A description of his instrument, as now in open within the Capitol at Washington, is all that I deem it pedient to introduce.

907. Upon a base board of wood, a U shaped bar of soil ma, w with covered copper wire, is firmly attached in a vertical posture. Q the terminal wires of the electro-magnet thus formed, communicate the distant station by means of a wire, the other by means of the pri A short distance above the electro-magnet, a har or keeper of sol \* attached at right angles to a lever. One extremity of this lever and steel point or tooth projecting downward, while through the other cut is a pivot supported by two stanchions behind the electro-magnet. At is attached to the upper surface of the lever tending to rame the, and it the keeper, from the poles of the electro-magnet beneath. When the cuit is completed with the voltaic battery at the distant station, the in magnet being charged, and suddenly attracting the keeper, cause in and the lever to descend. Upon the rupture of the circuit, attraction tween the magnet and keeper ceasing, the spring causes the lever to again, and thus, by repeated completions and interruptions of the co the lever may be made to play up and down, working upon the prof ( fulcrum. Immediately beneath the steel point a cylinder of wood, and with soft paper, is made to rotate upon an axis uniformly by means of a work. When, by the completion of the circuit, the steel point is made descend, it presses upon the soft paper of the revolving cylinder of long as the circuit remains unbroken. As the cylinder revolves, a dat: line of greater or less length is described upon the paper, and an aratr combination of such dots and lines constitutes the alphabet. Thus exletter may be written upon the paper at one extremity of a telegraph: of any length, by successive completions and interruptions of the circul the other.

908. As the velocity with which the electric current traverses a men wire has been found by Mr. Wheatstone to be about 289,000 mies a second, it is obvious that no perceptible interval of time will clapse a transmission of intelligence in this way, between any two points upon earth's surface.

909. Various modes of establishing the wires between two statess to been devised. It was at first proposed to enclose two insulated wires an iron pipe, buried beneath the surface of the earth; but this methodisectionable on the ground of expense, and liability to derangement above stated, Prof. Morse employs the earth itself for one-half of the cuand a wire, supported upon poles at some distance above the ground, a other.

### AN EFFORT

TO

### REFUTE THE ARGUMENTS ADVANCED

IN FAVOUR OF THE

## STENCE IN THE AMPHIDE SALTS, OF RADICALS,

CONSISTING,

LIKE CYANOGEN, OF MORE THAN ONE ELEMENT.

BY ROBERT HARE, M.D.
PROFESSOR OF CHEMISTRY IN THE UNIVERSITY OF PENNSYLVANIA.

### PHILADELPHIA:

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1842.



### AN EFFORT

To Refute the Arguments advanced in Favour of the Existence, in the Amphide Salts,\* of Radicals consisting, like Cyanogen, of more than one Element. By Robert Hare, M.D., Professor of Chemistry in the University of Pennsylvania.

The following is a Summary of the Opinions, which it is the Object of the subsequent Reasoning to Justify.

- (a) The community of effect, as respects the extrication of hydrogen by contact of certain metals with aqueous solutions of sulphuric and chlorohydric acid, is not an adequate ground for an inferred analogy of composition, since it must inevitably arise that any radical will, from any compound, displace any other radical, when the forces around its substitution preponderate over the quiescent affinities.
- (b) But if, nevertheless, it be held that the evolution of hydrogen from any combination, by contact with a metal, is a sufficient proof of the existence of a halogent body, simple or compound, in the combination, the evolution of hydrogen from water, by the contact with any metal of the alkalies, must prove oxygen to be a halogen body; also the evolution of hydrogen from sulphydric, selenhydric, or telluhydric acids, by similar means, would justify an inference that sulphur, selenium, or tellurium, as well as oxygen, belong to the halogen or salt radical class:—
- (c) The amphigen bodies being thus proved to belong to the halogen class, oxides, sulphides, selenides, and tellurides, would be haloid salts, and their compounds double salts, instead of consisting of a compound radical and a metal:—
- (d) The argument in favour of similarity of composition in the haloid and amphide salts, founded on a limited resemblance of properties in some instances, is more than counterbalanced by the extreme dissimilitude in many others:—
- (e) As, in either class, almost every property may be found which is observed in any chemical compound, the existence of a similitude, in some cases, might be naturally expected:—

are deemed salts, and which are consequently called haloid salts.

An amphide salt is one consisting of an acid and a base, each containing an amphigen body, either oxygen, sulphur, selenium, or tellurium, as its electro-negative ingredient.

The epithet halogen is applied to bodies whose binary compounds with metals

(f) As it is evident that many salts, perfectly analogous it position, are extremely dissimilar in properties, it is not a able to consider resemblance in properties, as a proof of small composition:—

(g) No line of distinction, as respects either properties of position, can be drawn between the binary compounds of a phigen and halogen bodies, which justifies that separate dution which the doctrine requires; so that it must be unteable

spects the one, or be extended to the other:-

(h) The great diversity, both as respects properties and position of the bodies called salts, rendering it impossible to the meaning of the word, any attempt to vary the larger theory of Chemistry, in reference to the idea of a salt, make tremely pernicious:—

(i) There is at least as much mystery in the fact, that! dition of an atom of oxygen to an oxacid, should confer an for a simple radical, as that the addition of an atom of this to such a radical, should create an affinity between it and an en

- (j) If one atom of oxygen confer upon the base into we enters, the power to combine with one atom of acid, it is quisistent that the affinity should be augmented, proportional further accession of oxygen:—
- (k) It were quite as anomalous, mysterious, and important there should be three expendentions, severally require saturation one, two, and three atoms of hydrogen, as that the meric states of phosphoric acid should exist, requiring as maderent equivalents of basic water:—
- (1) The attributes of acidity alleged to be due altogether presence of basic water, are not seen in hydrated acids, when ing water in that form only; nor in such as are, like the only incapable of uniting with water as a solvent. Further, these butes are admitted to belong to salts which, not holding wat base, cannot be hydrurets or hydracids of any salt radios while such attributes are found in compounds which, like the or carbonic acid, cannot be considered as hydrurets, they do set in all that merit this appellation, as is evident in the caprussic acid, or oil of bitter almonds:—
- (m) It seems to have escaped attention, that if SO be the sulphion of sulphates, SO, anhydrous sulphuric acid, must oxysulphion of the sulphites; and that there must, in the hyphites and hyposulphates, be two other oxysulphions!—

(n) The electrolytic experiments of Daniell have been

would so cause the separation of sodium, and oxygen, that the xygen would be attracted to the anode, the hydrogen and soda meing indirectly evolved by the reaction of sodium with water; while the acid, deprived of its alkaline base, would be found at the unode in combination with basic water, without having been made act in the capacity of an anion.

- (o) The copper in the case of a solution of the sulphate of this metal and a solution of potash, separated by a membrane, would, by electrolyzation, be evolved by the same process as sodium, so long as there should be copper to perform the office of a cathion; and when there should no longer be any copper to act in this capacity, the metal of the alkali, or hydrogen of water, on the other side of the membrane, would act as a cathion; the oxygen acting as an anion from one electrode to the other, first to the copper, and then to the potassium:—
- (p) The allegation that the copper was deposited from the want of an anion (oxysulphion) to combine with, is manifestly an error, since, had there been no anion, there could have been no discharge, as alleged, to hydrogen as a cathion, nor any electrolysis:—
- (q) The hydrated oxide precipitated on the membrane came from the reaction of the alkali with the sulphate of copper; the precipitated oxide of this metal from the oxygen of the soda acting as an anion; and the deposit of metallic copper from the solutions performing, feebly, the part of electrodes, while themselves the subjects of electrolyzation:—
- (r) The so called principles of Liebig,\* by which his theory of organic acids is preceded, are mainly an inversion of the truth, since they make the capacity of saturation of hydrated acids dependent on the quantity of hydrogen in their basic water, instead of making both the quantity of water, and, of course, the quantity of hydrogen therein, depend on their capacity:—
- (s) All that is truly said of hydrogen would be equally true of any other radical, while the language employed would lead the student to suppose that there is a peculiar association between capacity of saturation, and presence of hydrogen.

<sup>\*</sup> Traité de Chymie Organique, tom. 1, page 7.



## AN EFFORT,

&c. &c.

1. Some of the most distinguished European chemists, encouraged by the number of instances in which the existence of hypothetical radicals has meen rendered probable, have lately inferred the existence of a large numper of such radicals in a most important class of bodies heretofore consilered as compounds of acids and bases. It has been inferred, for instance, that sulphur, with four atoms of oxygen (SO4) constitutes a compound radizal, which performs in hydrous sulphuric acid, the same part as chlorine

in chlorohydric acid.

2. Graham has proposed sulphatoxygen as a name for this radical, and sulphatoxide for any of its compounds. Daniell has proposed oxysulphion and oxysulphionide for the same purposes. I have given the preference to a nomenclature moulded upon the last mentioned plan. In reasoning on the subject I shall use this nomenclature, not, however, with a view to manction it, as I disapprove altogether of this innovation, and deny the sufficiency of the grounds upon which it has been justified. Consistently with the language suggested by Daniell, hydrous sulphuric acid, constituted of one atom of acid and one of basic water, (SO3+HO) is a compound of oxysulphion and hydrogen (SO+H). Nitric acid (NO5+HO) is a compound of oxynitrion and hydrogen (NO6+H). In like manner we should have oxyphosphion in phosphoric acid, oxyarsenion in arsenic acid, and in all acids, hitherto called hydrated, whether organic or inorganic, we should have radicals designated by names made after the same plan. Their salts having corresponding appellations, would be oxysulphionides, oxynitrionides, &c. Also, in any salt in which any other of the amphigen class of Berzelius is the electro-negative ingredient, whether sulphur, selenium, or tellurium, all the ingredients excepting the electro-positive radical, would be considered as constituting a compound electronegative radical.\*

\* The conception of the existence of salt radicals seems to have originated with Davy. It was suggested by Berzelius, in his letter in reply to some strictures which I published on his Nomenclature, in the following language:—

"If, for instance, the true electro-chemical composition of the sulphate of potash should not be KO+SO3, as is generally supposed, but K+SO4, and it appears very natural that atoms, so eminently electro-negative as sulphur and oxygen, should be associated, we have, in the salt in question, potassium combined with a compound body, which, like cyanogen in K+C2 N, imitates simple halogen bodies, and gives a salt with potassium and other metals. The hydrated oxacids, agreeably to this view, would be then hydracids of a compound halogen body, from which metals may displace hydrogen, as in the hydracids of simple halogen bodies. Thus we know that SO3, that is to say, anhydrous sulphuric acid, is a body, whose properties, as respects acidity, differ from those which we should expect in the active principle of hydrous sulphuric acid.

"The difference between the oxysalts and the halosalts is very easily illustrated

3. It may be expedient to take this opportunity of mentioning, that is advocates of this new view, disadvantageously, as I think, employ the radical, to designate the electro-negative, as well as the electro-negative, as well as the electro-negative a compound halogen body. Cyanogen being analogous, is by him part in the halogen class. I shall, therefore, in speaking of "salt radical" improperly so called, employ the appellation contrived by the great Swant chemist.

4. It seems, however, to be conceded, that however plausible my to reasons for inferring the existence of halogen bodies in the amphic it would be inexpedient to make a corresponding change in nonceive on account of the great inconvenience which must arise from the convenience.

quent change of names.

5. Under these circumstances, it may be well to consider how far are is any necessity for adopting hypothetical views, to which it would be disadvantageous to accommodate the received language of chemists the strictures on the Berzelian nomenclature, which drew from Berzelian the suggestions contained in the quotation at the foot of the persent page, I stated it to be my impression that water should be considered acting in some cases as an oxybase, in others as an oxacid; and, is are amination of his reply,\* I observed that hydrous sulphuric acid with considered as a sulphate of hydrogen, and that when this coil with zinc or iron, the proneness of hydrogen to the aëriform and with zinc or iron, the proneness of hydrogen to the established land affinity.

Graham, Gregory, and myself, as respects the electro-positive related hydrogen to the amphigen and halogen elements, which I have designed collectively as the basacigen class; also in the impression that hydrogen acts like a metallic radical, its oxide, water, performing the part of a last I agree perfectly with Gregory in considering that hydrated acts allege that "acids and salts." But when the learned editor is allege that "acids and salts, as respects their constitution, will it are class," I consider him, and those who sanction this allegation, is an error upon an oversight. Because the salts of hydrogen, as the have water for their base, have heretofore been erroneously can be a salt of hydrogen.

one wrong name, cause all others to conform thereto!

'Silliman's Journal, Vol. 27, for 1835, page 61.

7. I fully concur with Gregory and Kane, in considering that were hydrous sulphuric acid, in nitric acid, chloric acid, and in organ is a generally acts as a base; also, that in this basic water hydrogen is a part perfectly analogous to that of a metallic radical: but, again is a

we are henceforth to confound salts with acids, and, instead of a restance

by formulæ. In KFF (fluoride of potassium), there is but one single has been tution, that is to say, that of K|FF; whilst in KOOOOS (sulphate of potassium) are two, K|OOOOS and KO|OOOS, of which we use the first in replaced metal by another, for instance, copper by iron; and the second in replacing are ide by another.

<sup>&</sup>quot;I do not know what value you may attach to this development of the costition of the oxysalts (which applies equally to the sulphosalts and others in the myself, I have a thorough conviction that there is therein something may be vague speculation, since it unfolds to us an internal analogy in phenomena. We agreeably to the perception of our senses, are extremely analogous

his view, I cannot perceive any difficulty in accounting for the evolution of nydrogen, as suggested in the quotation above made (6), agreeably to which, when diluted sulphuric acid reacts with zinc or iron, the liberation of hydrogen results from the superiority of the forces which tend to insert in the place occupied by the hydrogen, over those which tend to retain it in statu quo.

that the hydrogen unites with oxygen, and the chlorine with the metal; and hence it seems to be presumed, that when oxide of copper is combined with sulphuric acid, a similar play of affinities should ensue: but would it be reasonable to make this a ground for assuming the existence of a compound radical, when the phenomena admit of another explanation quite as simple and consistent with the laws of chemical affinity?

9. Whether hydrogen be replaced by zinc, or oxide of hydrogen by oxide of copper, cannot make any material difference. In the one case, a radical expels another radical, and takes its place; in the other, a base ex-

pels another base, and takes its place.

10. There can be no difficulty, then, in understanding wherefore, from the compound of sulphur and three atoms of oxygen, and an atom of basic water, hydrogen should be expelled and replaced by zinc, or that water should be expelled and replaced by oxide of copper; the only mystery is in the fact, that SO<sup>3</sup>, as anhydrous sulphuric acid, will not combine with hydrogen, copper, or any other radical, unless oxydized. But this mystery equally exists on assuming that an additional atom of oxygen converts SO<sup>3</sup> into oxysulphion, endowed with an energetic affinity for metallic radicals, to which SO<sup>3</sup> is quite indifferent.

11. In either case an inexplicable mystery exists; but it is, in the one case, associated with an hypothetical change, in the other, with one which

is known to take place.

12. But if hydrous sulphuric acid is to be assumed to be a hydruret of a compound halogen body (oxysulphion), because it evolves hydrogen on contact with zinc, wherefore is not water, which evolves hydrogen on contact with potassium, sodium, barium, strontium, or calcium, to be considered as a hydruret of oxygen, making oxygen a halogen body?

13. Boldly begging the question, Graham reasons thus: "the chlorides

themselves being salts, their compounds must be double salts."

14. But if the chlorides are salts, the chloride of hydrogen is a salt; and if so, wherefore is not the oxide of hydrogen a salt, which, in its susceptibility of the crystalline form, has a salt attribute which the aëriform chlo-

ride does not possess?

- 15. Further, if the oxide of hydrogen be a salt, every oxide is a salt, as well as every chloride. Now, controverting the argument above quoted, by analogous reasoning, it may be said, "the oxides themselves being salts, their compounds are double salts." Of course sulphate of potash is not a sulphatoxide, as Graham's ingenious nomenclature would make it, but must be a double salt, since it consists of two oxides in "themselves salts."
- 16. I trust that sufficient reasons have been adduced, to make it evident that the common result of the extrication of hydrogen, during the reaction of zinc or iron with sulphuric or chlorohydric acid, is not a competent ground for assuming that there are, in amphide salts, "compound radicals" playing the same part as halogen bodies.

17. Let us, in the next place, consider the argument in favour of the ex-

istence of such radicals, founded on the similitude of the baled and salts, which is stated by Dr. Kane in the following words:—

"It had long been remarked as curious, that bodies so different in position as the compound of chlorine with a metal, on one hand with exygen acid with the exide of the metal on the other, should be sed in properties that both must be classed as salts, and should not series of basic and acid compounds for the most part completely publications, p. 681.

18. Upon the similitude and complete parallelum of the mobile haloid salts, thus erroneously nileged, the author processis to arguing your of the existence in the former, of compound halogen bedee, unit

in their mode of combination to chlorine or jodine.

19. I presume it will be granted, that if simulatede in property but ficient ground for inferring an analogy in composition, desample to justify an opposite inference. And that if, as the author argue, of bodies have been classed as salts, on account of their similarity at spect, when dissimilar they ought not to be so classed. Under un the question, I propose to examine how far any similarity in proposition is to be be designated as salts by the author or any of mist.

20. The salts, hitherto considered as compounds of acids and have by Berzelius called amphide salts, being produced severally british with one or other of his amphigen class, comprising oxygen, while lenium, and tellurium, with two radiculs, with one of which we are od, with the other a base. The binary compounds of his taken comprising chlorine, bromine, iodine, fluorine and cyanogen, as of the compounds.

him haloid salts. I shall use the names thus suggested.

21. Among the haloid salts we have common salt and Drophing the gaseous fluorides and chlorides of hydrogen, silicon or have: the ing liquor of Libavius; the acrid butyraceous chlorides of the animony; the volatile chlorides of magnesium, iron, chroman, and cury, and the fixed chlorides of calcium, barriam, strontiam, which lead; the volatile poison prussic acid, and solid poisonous because cury, with various inert cyanides like those of Prussian blue; alternated great number of etherial compounds.

22. Among the amphide salts are the very soluble sulphates of 1 iron, copper, soda, magnesia, &c., and the insoluble stony sulphates of ryta and strontia; also ceruse and sugar of lead; alabaster, markle, at ethers, and innumerable stony silicates, and aluminates. Last, but among the least discordant, are the hydrated acids, and alkaline and on

hydrates.

28. When the various sets of bodies, above enumerated, as compute the two classes under consideration, are contemplated, is it not ended not only between several sets of haloid and amphide salts, but also is several sets in either class, there is an extreme discordancy in proper so that making properties the test would involve not only that varies in one class could not be coupled with certain sets in the other, but that in neither class could any one set be selected as exemplifying the racteristics of a salt, without depriving a majority of those similarly of tuted, of all pretensions to the saline character?

24. Now, if among the bodies above enumerated, some pairs of and and haloid salts can be selected, which make a tolerable match with re

• their properties, as in the case of sulphate of soda, and chloride of sodium, hile in other cases there is the greatest discordancy, (as in the stony silithe felspar, and the gaseous fluoride fluosilicic acid gas; as in soap and erbyshire spar; as in marble and the fuming liquor of Libavius, the sour potochloride of tin, and sweet acetate of lead), is it reasonable to found an rgument in favour of a hypothetical similitude in composition, on the reemblance of the two classes in properties? Does not the extreme dissimitude in some cases, more than countervail the limited resemblance in thers? And when the great variety of properties displayed both by the amphide and haloid salts is considered, is it a cause for wonder or perplexity, rat in some instances, amphide salts should be found to resemble those of **28 other kind?** 

25. Again, admitting that there was any cause for perplexity agreeably > the old doctrine, is there less, agreeably to that which is now recommendd? Is there no ground for wonder that oxygen or sulphur cannot act as imple halogen bodies? By what rule are their binary compounds to be **xcluded** from the class of haloid salts? Wherefore should chlorides, bronides, iodides, and fluorides, however antisaline in their properties, be conidered as salts, while in no case is an oxide, a sulphide, selenide or tellu-

ide to be deemed worthy of that name?

26. I challenge any chemist to assign any good reason wherefore the red odide of mercury is any more a salt than the red oxide, or the protochloide is more saline than the sulphide: or why the volatile oxides of osmium of arsenic are less saline than horn silver or horn lead; or the volatile **phloride** of arsenic, than the comparatively fixed sulphides of the same metal:

why gaseous chlorohydric acid is more saline than steam or gaseous oxsydric acid.

27. It much surprises me, that when so much stress is laid upon the idea of a salt, the impossibility of defining the meaning of the word escapes atention. How is a salt to be distinguished from any other binary com-When the discordant group of substances which have been enumerated under this name is contemplated, is it not evident that no definition of them can be founded on community of properties; and, by the advocates of the new doctrine, composition has been made the object of definition, instead of being the basis; thus, agreeably to them, a compound is not a salt, because it is made of certain elements; but, on the contrary, an element, whether simple or compound, belongs to the class of salt radicals, because it produces a salt. Since sulphur, with four atoms of oxygen, SO4, produces a salt with a metal, it must be deemed a salt radical.

28. In proof that the double chlorides are not united in a way to justify the opinion adopted by Bonsdorff, Thomson, myself, and others, it is alleged by Graham, "that in such compounds the characters of the con-

statuent salts are very little affected by their state of union."

- 29. This allegation being, in the next page, admitted to be inapplicable in the case of the double cyanides; an effort is made to get over this obstacle, by suggesting the existence of another compound radical. But the allegation of the author is erroneous as respects various double haloid salts, especially the fluosilicates, the fluoborates, fluozirconiates, the chloroplatipates, chloroiridiates, chloroosmiates, chloropalladiates, &c., all of them compounds in which the constituent fluorides and chlorides exist in a state of energetic combination by which they are materially altered as to their state of existence.
  - 30. Evidently the word salt has been so used, or rather so abused, that



gen, hydroxion, and these compounds would conceive that the haloid compounds, erroneou more correctly considered as single salts, can t

32. Between the reaction of fluoboric acid v acid with oxybases, is there not a great resemb

33. I am unable to understand how, if the exysalts is inferred, the other salts of the ampl from a corresponding inference. But if the ex. double sulphides be admitted, can it be consisalso in double chlorides, iodides, &c.? Is there tween the habitudes of sulphur, selenium, and those of the halogen bodies?

34. Would not the modification of the etheri the new hypothesis, be disadvantageous, both a ception of those compounds, and the names whice priate! Would not the transfer of the oxygen fi acid, and the creation, thus, of new salt radicals objectionable; such as oxyoxalion for oxalates. oxyacetion for acetates; while, for their compooxalionides, oxytartarionides, oxyacctionides, &

35. If sulphates are to be considered as oxys are we to designate the sulphites, hyposulphites So O3, So O3? SO3 may, perhaps, with more consisting of a compound radical, SOs, and o sulphurous acid; but in a sulphite, anhydrous s a species of oxysulphion itself, being as much t philes, as SO<sup>4</sup> is of the sulphates. Of course & finity for radicals, contrary to fact. I presume to be trioxysulphionides; hyposulphites, sesquiquadroxysulphionides; while the hyposulphates quintoxysulphionides!!!

36. Analogous complication in nomenclatur the nitrites and nitrates, phosphites and phosp ates; also as respects the carbonic, and oxalic a

37. It is true that nature has not so made her ennarated into obsesses between which any distir

creating the compounds formed by electro-negative elements with anions, cas acids; those formed with cathions, as bases; while the combinations corned by the union of such acids and bases have been considered as simse salts. Thus four classes are constituted, consisting of electro-negative relements, of acids, bases, and single salts, while, by the union of the later, a fifth class of double salts is formed. Whether the words acid, base, and salt, be adhered to, objectionable as they are in some respects, and esexecially the latter, or some others be contrived, it would seem to me disadvantageous to merge them in one name, pursuant to the views of the advosates of salt radicals, as stated by Gregory in his edition of Turner's Chemistry, 572.

38. The objection, that not being electrolytes the relation of acids and bases to the voltaic electrodes cannot be discovered, is easily remedied; since, on the union of a common ingredient with an anion and a cathion, there cannot be any doubt that the resulting compounds will have the same electro-chemical relation as their respective heterogeneous ingredients; so that, with the anion, an acid or electro-negative body will be formed; with the cathion, a base or electro-positive body. Moreover, as respects organic compounds which cannot be subjected to the electrolytic test, whatever saturates an inorganic acid must be a base, and whatever saturates an inor-

ganic base must be an acid.

39. The word salt, I have shown, is almost destitute of utility, from the impossibility of defining it, and the amplitude of its meaning. means every thing, is nearly as useless as that which means nothing.

40. As respects the three phosphates of water, PO5+HO, PO5+2HO, PO+3HO, the argument used by Dr. Kane cuts both ways; although, by its employer, only that edge is noticed which suits his own purpose. alleged that the difference of properties, in these phosphates, is totally inexplicable upon the idea of three degrees of "hydration;" but that all difficulty vanishes, when they are considered as three different compound salt radicals, oxyphosphionides of hydrogen, PO<sup>6</sup>+H, PO<sup>7</sup>+2H, PO<sup>8</sup>+3H.

41. To me the formation of three compound elements, by the reiterated addition of an atom, of which five of the same kind were previously in the mass to which the addition is made, seems more anomalous, mysterious, and improbable, than the existence of three compounds of phosphoric acid with water, in which the presence of the different proportions of water is the consequence of some change in the constitution of the elements which is referred to isomerism.

42. No reason can be given why the addition of one, two, and three atoms of oxygen, to the "radical," should convey a power to hold a proportional number of atoms of hydrogen. Such an acquisition of power is an anomaly.

43. In the case of radicals formed with hydrogen in different proportions, as in acetyl and ethyl, formyl and methyl, the number of atoms of oxygen in the peroxides, is the inverse of the hydrogen in the radical.

44. Ethyl, C4, H5, unites, at most, with one atom of oxygen, while acetyle, C4, H2, takes three atoms to form acetic acid, C4, H2, O2. Methyl, C<sup>2</sup>, H<sup>2</sup>, forms, in like manner, only a protoxide, while formyl, C<sup>2</sup>, H, takes three atoms to constitute formic acid.

45. Besides the three oxyphosphions, of which the formulas are above stated, there would have to be another in the phosphites; so that instead of the hydrated acid, or phosphite of water, being PO<sup>2</sup>+HO, it would have to be PO+H, a fourth oxyphosphionide of hydrogen.

46. Respecting the new principles which I have been contents, it Kane alleges "that the elegance and simplicity with which the laws of so line combination may be traced from them is remarkable," because keep ceives, that without an appeal to those principles, the fact that the unter of equivalents of acid in a salt are proportionable to the number of expelents of oxygen in the base, would be inexplicable.

47. Thus, when the base is a protoxide, we have one atom of the provide of hydrogen to take its place; when the base is a resquande to of radical and three of oxygen), three atoms of the protoxide of hydrogen take its place: if the base be a bioxide, two atoms of the protoxide of by-

drogen take its place.

48. I have already adverted to the existence of certain chemical and inexplicable in the present state of human knowledge. Among there that of the necessity of oxidation to enable metallic radicals to comme with acids. But as a similar mystery exists as respects the adventure property of combining with radicals, which results from the acquisitor of an additional atom of oxygen by any of the compounds hitherto consider as anhydrous acids, the new doctrine has in that respect no processes claim to credence.

49. But if, without impairing the comparative pretensions of the presing doctrine, we may appeal to the fact that the acquisition of an atom of an atom of an atom of acid, is not consistent that the acquisition of two atoms of oxygen should consistent to hold two atoms of acid, and that with each further acquisitor of oxygen a further power to hold acids should be conferred?

50. So far then there is in the old doctrine no more inscrutability has in that which has been proposed as its successor. Since if on the one has it be requisite that for each atom of oxygen in the base, there shall be as atom of acid in any salt which it may form, on the other, in the case of the three oxyphosphions, for each additional atom of hydrogen extrancous transcriptions and the results and the results and the results are salt radical, there must be an atom of oxygen superadded to the results.

- 51. It being then admitted that, numerically, the atoms of exponents oxysalt will be as the atoms of oxygen in the base, it must be a substant whenever an oxysalt of a protoxide is decomposed by a bookle, were have to be two atoms of the former for one of the latter. For the has two atoms of oxygen, and requires by the premises two atoms while the salt of the protoxide, having but one atom of oxygen, and yield, only one atom of acid. Two atoms of this salt, the rest is there its base be water, or any other protoxide, will be decome atom of bioxide; provided the affinity of the acid for the look is atoms of the acid for the look.
- 52. It follows, that the displacement of water from its such is the by Kane, does not favour the idea that hydrous suiphure action vesulphionide of hydrogen, more than the impression that it is is the water.

53. Of course, in the case of presenting either a sesquex by cravity ide, to the last mentioned sulphate, in other words, hydrous salignate, the same rationale will be applicable.

54. The next argument advanced by Dr. Kane, is, that some of the of which the existence is assumed upon the old doctrine, and layer as they have never been isolated. This mode of reasoning this to react against the new doctrine with pre-eminent force, since all its

compound radicals imagined by it are hypothetical—none of them having been isolated.

- 55. The third argument of the respectable author above named is, that acids display their acid character in a high degree only when in the combination with water.
- 56. This argument should be considered in reference to two different cases, in one of which all the water held by the acid is in the state of a base, while in the other an additional quantity is present acting as a solvent. So far as water, acting as a solvent, facilitates the reaction between acids and bases, it performs a part in common with alcohol, ether, volatile oils, resins, vitrifiable fluxes, and caloric. Its efficacy must be referred to the general law, that fluidity is necessary to chemical reaction. "Corpora non agunt nisi soluta."
- 57. In a majority of cases, basic water, unaided by an additional portion acting as a solvent, is quite incompetent to produce reaction between acids and other bodies. Neither between sulphuric acid and zinc, between nitric acid and silver, nor between glacial or crystallized acids and metallic oxides, does any reaction take place without the aid of water acting as a solvent, and performing a part analogous to that which heat performs in promoting the union of those oxybases with boric, or silicic acid.

55. It is only with soluble acids that water has any efficacy. The difference between the energy of sulphuric and silicic acid, under the different circumstances in which they can reciprocally displace each other, is founded on the nature of the solvents which they require, the one being only capable

of liquefaction by water, the other by caloric.

- 59. In support of his opinions the author adverts to the fact, that with bydrated sulphuric acid, baryta will combine energetically in the cold, while a similar union between the anhydrous vapour and the same base cannot be accomplished without heat. But it ought to be recollected, that to make this argument good, it should be shown wherefore heat causes the baryta, a perfectly fixed body, to unite more readily with an aëriform substance in which increase of temperature must, by rarefaction, diminish the number of its particles in contact with the solid. If the only answer be, that heat effects some mysterious changes in affinity, (or as I would say in the electrical state of the particles) it should be shown that the presence of water or any other base has not been productive of a similar change, before another explanation is held to be necessary. But I would also call to mind that the hydrated acid is presented in the liquid state; and if it be asked why water, having less affinity than baryta, can better cause the condensation of the acid, I reply, that it is brought into contact with the acid both as a liquid and a vapour, of neither of which forms is the earthy base susceptible. But if all that is necessary to convert anhydrous sulphuric acid into an oxysulphionide, be an atom of oxygen and an atom of metal, what is to prevent baryta and anhydrous sulphuric acid from forming an oxysulphionide of barium? All the elements are present which are necessary to form either a sulphate or oxysulphionide; and I am unable to conceive wherefore the inability to combine does not operate as much against the existence of radicals as of bases.
- 60. I would be glad to learn why, agreeably to the salt radical theory, anhydrous sulphuric acid unites with water more greedily than with baryta, and yet abandons the water promptly on being presented to this base. Why should it form an oxysulphionide with hydrogen more readily than with barium, and yet display, subsequently, a vastly superior affinity for barium?

61. It seems to be overlooked, that anhydrous sulphuric and him oxysulphion of the sulphites, ought to form sulphites on contact with

62. But if the sulphate of water owe its energy to that pormitiond, which, by its decomposition gives rise to the compount able sulphion, and not to the portion which operates as a solvent, what the concentrated state, will it not react with iron and zinc, without water, when, with dilution, it reacts most powerfully with those

63. Some stress has been laid upon the fact, that sources a tecived, excepting with the aid of water, as if to derive force for the doctrine from that old and popular, though now abundanced text of the but it should be recollected that it is not the water which goes to tecompound element in the "hydracids," erroneously so called, this fers sourcess. Will any one pretend that either sulphure or the when concentrated, is sour? Are they not caustic! Car are crystallized organic acids be said to have a sour tuste, independent moisture of the tongue? The hydrated oily acids being incapability with water as a solvent, have none of these vulgar attributes of the absence of these attributes in prussic acid would alone be will render it inconsistent to consider them as having any connexional presence of hydrogen.

with exides on contact. To this I would add, that it does not exact with exides on contact. To this I would add, that it does not exact water under those circumstances, but, on the contrary, specially like oil, after mechanical mixture; nor does it, under any circum unite with an equivalent proportion of water to form a hydrar. Of as it is not to basic water that it is indebted for its ability to become gredient in salts, it cannot be held that this faculty is the reset of

vious conversion into an oxycarbionide of hydrogen.

not, therefore, be considered as exychromionide of hydrogen. Let of bitter almonds, which consists of a compound radical, beautiful atom of hydrogen, and which is therefore constituted precisit at radical doctrine requires for endowment with the attributes of an "hidi is utterly destitute of that acid reaction which hydrogen is repost peculiarly competent to impart. It follows that we have, on the or in chromic acid, a compound endowed with the attributes of acid; out being a hydroret of any compound radical; and, on the other a bitter almonds, a hydroret of a compound radical, without any of he butes of acidity.

66. The last argument in favour of the existence of salt radicals. I have to answer, is that founded on certain results of the electric

saline solutions.\*

Faraday found that when various substances were electrolyzed, a volument at the same time in the circuit, that for every equivalent of water decomposed the tube, neither more nor less than an equivalent of the other body could be a

posed.

<sup>\*</sup> It is well known that Faraday employed a very simple instrument to at the quantity of the gaseous elements of water yielded in a given time insubjected to the voltaic current. It consisted of a graduated tube through set of which the current was conveyed by wires, so terminating within it is to interval between them through which the current, being conveyed by the city process, effected the decomposition of the intervening liquid, the results being caught and measured by the tube. This instrument has been called a electrometer, or voltameter

sexposed to the current employed, simultaneously with some water in a volumeter, Daniell alleges that, for each equivalent of the gaseous elements of water evolved in the voltameter, there was evolved at the cathode and anode, not only a like quantity of those elements, but likewise an equal number of equivalents of soda and sulphuric acid. This he considers as involving the necessity, agreeably to the old doctrine, of the simultaneous decomposition of two electrolytic atoms in the solution, for one in the voltameter; while, if the solution be considered as holding oxysulphionide of sodium, instead of sulphate of soda, the result may be explained consistently with the law ascertained by Faraday. In that case, oxysulphion would be carried to the anode, where, combining with hydrogen, it would cause oxygen to be extricated, while sodium, carried to the cathode, and deoxidizing water, would cause the extrication of hydrogen.

68. Dr. Kane, alluding to the experiments above mentioned, and some others which I shall mention, alleges that "Professor Daniell considers

the binary theory of salts to be fully established by them."

69. Notwithstanding the deference which I have for the distinguished inventor of the constant battery, and disinclination for the unpleasant task of striving to prove a friend to be in the wrong, being of opinion that these inferences are erroneous, I feel it to be my duty, as a teacher of the science, to show that they are founded upon a misinterpretation of the facts appealed to for their justification.

70. It appears to me, that the simultaneous appearance of the elements of water, and of acid and alkali, at the electrodes, as above stated, may be accounted for, simply by that electrolyzation of the soda, which must be the natural consequence of the exposure of the sulphate of that base in the circuit. I will, in support of the exposition which I am about to make, quote the language of Professor Daniell, in his late work, entitled, "Introduction to Chemical Philosophy," page 413:—

"Thus we may conceive that the force of affinity receives an impulse which enables the hydrogen of the first particle of water, which undergoes decomposition, to combine momentarily with the oxygen of the next particle in succession; the hydrogen of this again, with the oxygen of the next; and so on till the last particle of hydrogen communicates its impulse to the platinum, and escapes in its own elastic form."

71. The process here represented as taking place in the instance of the oxide of hydrogen, takes place, of course, in that of any other electrolyte.

72. It is well known, that when a fixed alkaline solution is subjected to the voltaic current, that the alkali, whether soda or potassa, is decomposed; so that if mercury be used for the cathode, the nascent metal, being protected by uniting therewith, an amalgam is formed. If the cathode be of platinum, the metal, being unprotected, is, by decomposing water, reconverted into an oxide as soon as evolved. This shows, that when a salt of potassa or soda is subjected to the voltaic current, it is the alkali which is the primary object of attack, the decomposition of the water being a secondary result.

73. If in a row of the atoms of soda, extending from one electrode to the other, while forming the base of a sulphate, a series of electrolytic decompositions be induced from the cathode on the right, to the anode on the left, by which each atom of sodium in the row will be transferred from the atom of acid with which it was previously combined, to that next upon the right,

causing an atom of the metal to be liberated at the cathode; this see oxidizing water, will account for the soda and hydrogen at the a Meanwhile the atom of sulphate on the left, which has been depressed sodium, must simultaneously have yielded to the anode the oxygen by this metal was oxidized. Of course the acid is left in the hydrou usually called free, though more correctly esteemed to be that of a see of water.

74. I cannot conceive how any other result could be expected it electrolysis of the base of sulphate of soda, than that which is a scribed. Should any additional illustration be requisite, it will be a a note subjoined.\*

75. I will, in the next place, consider the phenomena observed: fessor Daniell, when solutions of potassa and sulphate of copper, so by a membrane, were made the medium of a voltaic current.

76. Of these I here quote his own account. Philosophical Magaz

Journal, Vol. 17, p. 172:—

"A small glass bell, with an aperture at top, had its mouth close ing a piece of thin membrane over it. It was half filled with a dilution of caustic potassa, and suspended in a glass vessel containing a

\* It is easy to understand how a simultaneous appearance of oxygen and the anode, and soda and hydrogen at the cathode, may ensue, simply by the lyzation of the alkaline base from the following association of formulæ.

Anhydrous sulphuric acid is represented by the usual formula, SO: on the usual symbol, O; sodium by Na; water, acting as a solvent, by HO E of oxygen, sodium, or acid, is numbered from right to left, 1, 2, 3, 4, so change of position consequent to electrolysis may be seen.

| Anode | 1<br>O                     |                                   | 3<br>O                     |                                                                     | 4 Water<br>O HO Cal |      |
|-------|----------------------------|-----------------------------------|----------------------------|---------------------------------------------------------------------|---------------------|------|
|       | Na Na                      | Na Na                             |                            | $\widetilde{N_a}$                                                   | <sup>4</sup><br>Na  |      |
|       | 1<br>SO <sup>3</sup><br>HO | 2<br>~~~<br>SO <sup>3</sup><br>HO |                            | 3<br>SO <sup>3</sup><br>HO                                          | 30°<br>HO           |      |
| Anode | 1 ()                       | O<br>Na                           | 3<br>O<br>2<br>Na          | $ \begin{array}{c} \frac{4}{O} \\ 3 \\ \widetilde{Na} \end{array} $ | O<br>4<br>Na        | H '- |
|       | SO <sup>3</sup><br>HO      | 2<br>SO3<br>HO                    | 3<br>SO <sup>3</sup><br>HO | 4<br>SO <sup>3</sup><br>HO                                          |                     |      |

As the atoms are situated in the second arrangement, the atom of at 2"

at the anode, the atom of sodium, Na, with which it had been under transferred to the second atom of sulphuric acid, which had yielded as seed

third atom of acid, SO3, this having, in like manner, yielded its sodium to to

atom of acid, SO3, from which the fourth atom of sodium. No. had beer 25 by the electrolytic power. The atom of sodium thus removed from the four of acid, is represented in union with the oxygen of an atom of water, of hydrogen, H, is at the cathode.

reutral solution of sulphate of copper, below the surface of which it just lipped. A platinum electrode, connected with the last zinc rod of a large mentant battery of twenty cells, was placed in the solution of potassa; and mental copper immediately under the diaphragm which separated the two solutions. The circuit conducted very readily, and the action was very entretic. Hydrogen was given off at the platinode in a solution of potassa, and oxygen at the zincode in the sulphate of copper. A small quantity of gas was also seen to rise from the surface of the diaphragm. In about ten minutes the lower surface of the membrane was found beautifully coated with metallic copper, interspersed with oxide of copper of a black colour, and hydrated oxide of copper of a light blue.

cell we have two electrolytes separated by a membrane, through both of which the current must pass to complete its circuit. The sulphate of copper is resolved into its compound anion, sulphuric acid + oxygen (oxysulphion), and its simple cathion, copper: the oxygen of the former escapes at the zincode, but the copper on its passage to the platinode is stopped at the surface of the second electrolyte, which for the present we may regard as water improved in its conducting power by potassa. The metal here finds nothing by combining with which it can complete its course, but being forced to stop, yields up its charge to the hydrogen of the second electrolyte,

which passes on to the platinode, and is evolved.

"The corresponding oxygen stops also at the diaphragm, giving up its charge to the anion of the sulphate of copper. The copper and oxygen thus meeting at the intermediate point, partly enter into combination, and form the black oxide; but from the rapidity of the action, there is not time for the whole to combine, and a portion of the copper remains in the metallic state, and a portion of the gaseous oxygen escapes. The precipitation of blue hydrated oxide doubtless arose from the mixing of a small portion of the two solutions."

Faraday, there are two modes in which a voltaic current may be transmitted, conduction and electrolyzation. In order that it may pass by the last mentioned process, there must be a row of anions and cathions forming a series of electrolytic atoms extending from the cathode to the anode. It is not necessary that these atoms should belong to the same fluid. A succession of atoms, whether homogeneous, or of two kinds, will answer, provided either be susceptible of electrolyzation. Both of the liquids resorted to by Daniell, contained atoms susceptible of being electrolyzed. If his idea of the composition of sulphate of copper, and the part performed by the potassa, were admitted for the purpose of illustration, we should, on one side of the membrane, have a row of atoms consisting of oxysulphion and copper; on the other, of oxygen and hydrogen.

78. Recurring to Daniell's own description of the electrolyzing process, above quoted, an atom of copper near the anode being liberated from its anion, oxysulphion, and charged with electricity, seizes the next atom of oxysulphion, displacing and charging an atom of copper therewith united. The cupreous atom thus charged and displaced, seizes a third atom of oxysulphion, subjecting the copper, united with it, to the same treatment as it had itself previously met with. This process being repeated by a succession of similar decompositions and recompositions, an electrified atom of copper is evolved at the membrane, where there is no atom of oxysulphion. Were there no other anion to receive the copper, evidently the electrolyzation would not have taken place; but oxygen, on the one side of the membrane, must succeed to the office performed by oxysulphion on the other



pagation of that electrolytic process, by which as alleged, "pass to the platinode," i. e. catho 80. In these strictures I am fully justified by Faraday, which I quote from his Researches, #

"A single ion, i. e. one not in combination widency to pass to either of the electrodes, and withe passing current, unless it be itself a compound so subject to actual decomposition."

"If, therefore, an ion pass towards one of must also be passing simultaneously to the othe secondary action, it may not make its appearat

81. In explanation of the mixed precipitate brane, I suggest that the hydrated oxide resul between the alkah and acid, the oxide from the assa acting as a cathion in place of that of the o metallic copper is to be attributed to the solution and as electrolytes; so that, at the membrane formed, which enabled a portion of the coppe combining with an anion, and a portion of oxyguniting with a cathion. In this explanation I a account of a well known experiment by Farac magnesia and water was made to act as elect spectively.

82. There can, I think, be no better proof placed on the experiments with membranes, in the existence of compound radicals in acids is 1 into which an investigator, so sagacious as my been led, in explaining the complicated results.

83. The association of two electrolytes, and tween the potassa and acid, which is admitted to oxide, seem rather to have created difficulties th

84. In this view of the subject, I am support day, as expressed in the following language:—

"When other metallic solutions are used, co exides, as that of copper combined with this or 85. I cannot conceive, that in any point of view the complicated and "confused" results of the experiment of Daniell with electrolytes separated by membranes, are rendered more intelligible by supposing the existence of salt radicals. I cannot perceive that the idea that the anion in the sulphate is oxysulphion, makes the explanation more satisfactory than if we suppose it to be oxygen. Were a solution of copper subjected to electrolysis alone, if the oxide of copper were the primary object of the current, the result would be analogous to the case of sodium, excepting that the metal evolved at the cathode, not decomposing water, would appear in the metallic form. If water be the primary object of attack, the evolution of copper would be a secondary effect.

86. It is remarkable, that after I had written the preceding interpretation of Daniell's experiments, I met with the following deductions stated by Matteuchi, as the result of an arduous series of experiments, without any reference to those of Daniell above mentioned. It will be perceived that these

deductions coincide perfectly with mine.

87. I subjoin a literal translation of the language of Matteuchi from the Annales de Chimie et de Physique, tome 74, 1840, page 110:—

"When salt, dissolved in water, is decomposed by the voltaic current, if the action of the current be confined to the salt, for each equivalent of water decomposed in the voltameter, there will be an equivalent of metal at the negative pole, and an equivalent of acid, plus an equivalent of oxygen, at the positive pole. The metal separated at the negative pole will be in the metallic state, or oxidized according to its nature. If oxidized, an equivalent of hydrogen will be simultaneously disengaged by the chemical decomposition of water."

88. Thus it seems, that the appearance of acid and oxygen at the anode, and of alkali and hydrogen at the cathode, which has been considered as requiring the simultaneous decomposition of two electrolytes upon the heretofore received theory of salts, has, by Matteuchi, been found to be a result requiring the electrolysis of the metallic base only, and, consequently, to be perfectly reconcilable with that theory.

89. In fact I had, from the study of Faraday's Researches, taken up the impression, that the separate appearance of an acid and base, previously forming a salt, at the voltaic electrodes, was to be viewed as a secondary effect of the decomposition of the water or the base; so that acids and bases

were never the direct objects of electrolytic transfer.

## Of Liebig's "Principles," so called.

- 90. Under the head of the "theory of organic acids," in Liebig's Treatise on Organic Chemistry, we find the following allegations dignified by the name of principles. Manifestly they must tend to convey a false impression to the student, that hydrogen has a peculiar property of creating a capacity for saturation, instead of being only the measure of that capacity, as is actually true, and likewise that in this respect it differs from any other radical.
- 91. The allegations to which I refer are as follows, being a literal translation from the French copy of the Traité of Liebig, page 7:—
- "The hydrated acids are combinations of one or more elements with hydrogen, in which the latter may be replaced wholly or in part by equivalents of metals."
- "The capacity of saturation depends consequently on the quantity of hydrogen which can be replaced.

"The compound formed by the other elements being comideral was dical, it is evident that the composition of this radical can exercise and ence on the capacity of saturation.

"The capacity of saturation of these achie augments or dimension in same ratio as the quantity of hydrogen, not entering into the 👊 👊

augments or diminishes.

"If into the composition of the salt radical there should be introduced undetermined quantity of any elements, without changing the carrie hydrogen extraneous to the radical, the atomic weight of the extra be augmented, but the capacity of saturation would remain the mac."

92. As by the advocates of the existence of "salt radicals," in the is considered as playing the part of a metallic radical, and most were as respects any relation between it and the capacity of anturates has same predicament as any other electro-positive radical, I canada and wherefore laws, which affect every other body of this kind, should a selfas if particularly associated with hydrogen.

93. Would not a more comprehensive and correct idea be promised

the following language?—

94. From any combination of an acid with a base, enter the lar ve radical may be replaced by any other radical or base, between west al the other elements present, there is a higher affinity. Of course from and called hydrated, from their holding an atom of basic water, either as 🗷 or its radical (hydrogen), may be replaced by any other compress the a radical.

95. The premises being manifestly fallacious, still more so to the state quent allegation, that in consequence of the hydrated acids less to pounds formed with hydrogen, their capacity of saturation deposits

quantity of this element which can be replaced.

96. Is not this an inversion of the obvious truth, that the same hydrogen present is as the capacity of saturation; and that, of ours, in quantity of any element which can be substituted for it, must be a quantity lent proportion? Would not a student, from this, take up two enumers

\* There is, in some respects a coincidence so remarkable as to the per an " Dr Kane and myself, with respect to hydrogen, that I quote here the

which has been he d by us respectively on this subject

Treating of hydrogen, Dr Kane uses the following words - It was the of the compounds of hydrogen combined to show that it was at epositive body that it to k place along with iron, manganese, and in These views have been still further corroborated by the research There rests now no doubt, in the minds of philosophica con the drogen is a matal enor neasty volatile.

To s justifies the following language held in my letter on the Bern it was

"I am of opinion that the employment of the word hydraeid is oxagid, must find the invery the errongous idea, with which, in present definition, the outhor sector to have been imbued, that had sight in the plays the same part as oxygen in the other. But in reality, the icraes a very a combustible, and of course the radical, by his own defin ton

So entirely have I can tirred in considering hydrogen as an air fire, city is for more than twenty years. I have, in my lectures, accounted for the same of mercury when e ectrolysed in contact with sal amm mixe, by mercing a sale to be a gaseous alloy of two metallic ingredients, hydrogen and introgen to it was

ačriform metals.

- —first, that the capacity of saturation is conferred by the radical, and a next place, that of all radicals, hydrogen alone can give such a cay? Is it not plain, that the assertion here made by the celebrated auwould be true of any radical?
- Passing over a sentence which has no bearing on the topic under ssion, in the fourth allegation we have a reiteration and expansion of rror of those by which it is preceded. We are informed that the "caty of saturation augments and diminishes with the quantity of hyen which can be replaced," which is again an inversion of the truth, the quantity of hydrogen varying with the capacity, the quantity of other radical, competent to replace it, must be in equivalent propor-
- Is not the concluding allegation a mere truism, by which we are med, "that if any undetermined quantity of any element should be inced into the composition of the radical, without changing the capacity neasured by hydrogen), the capacity would be found the same when ured by any other radical?"
- As all that is thus ascribed to hydrogen must be equally true of other radical, there would have been less liability to misapprehension, he generic term radical been employed wherever hydrogen is mend. But by employing the word radical to designate halogen elements, dvocates of the existence of compound radicals in amphide salts have ved the word in question of much of its discriminating efficacy. In their nomenclature would confound all ultimate elements under one ric appellation, and all their binary combinations under another, so that st every chemical reagent, whether simple or compound, would be a or a radical.
- 0. Before concluding, I feel it to be due to the celebrated German cheabove mentioned, to add, that however I may differ from him as to the being hydrurets of compound radicals, I am fully disposed to make owledgments for the light thrown by his analytical researches on orchemistry, and the successful effect of his ingenious theoretic specuas, in rendering that science more an object of study with physicians agriculturists.



### AN ABSTRACT

From Kane's Elements, of the Arguments in Favour of the Existence of Compound Radicals in Amphide Salts.

It appears proper to give the student of this text book, the option of studying the arguments which it is the object of the preceding pages to refute. Hence I subjoin the following abstract from Kane's Elements, page 681:—

"It had been long remarked as curious, that bodies so totally different in composition as the compound of chlorine with a metal on the one hand, and of an oxygen acid with the oxide of the metal on the other, should be so similar in properties, that soth must be classed together as salts, and should give origin to series of basic and acid compounds for the most part completely parallel. This difficulty has been so much felt by the most enlightened chemists, that doubts have been raised as to whether the acid and base, which are placed in contact to form by their union an oxygen salt, really exist in it when formed; and it has been suggested, that at the moment of union a new arrangement of elements takes place, by which the structure of the resulting salt is assimilated to that of a compound of chlorine or of iodine with a metal. This view, at first sight so far-fetched, which considers that in glauber's salt there is neither sulphuric acid, nor soda, but sulphur, oxygen, and sodium, in some other and simpler mode of combination, is now very extensively received by chemists; and I shall proceed, therefore, to describe with some detail the form which it has assumed, and the evidence by which it is supported.

The greater number of those bodies which are termed oxygen acids, have not been in reality insulated, and what are popularly so called are merely supposed to contain the dry acid combined with water. Thus the nearest approach we can make to nitric acid, is the liquid NO6H; to acetic acid, the crystalline body C4H4O4; and to oxalic acid, the sublimed crystals C2O4H; we look upon these bodies as being combinations of the dry acid with water, and we write their formulæ NO5+HO, and C4H3O3+HO and C2O3+HO, but that these dry acids exist at all is a mere assumption. Hence with regard to these instances, and they embrace the majority of all known acids, the idea that the acid and base really exist in the salt formed by the

action of hydrated acids on a base, is purely theoretical.

When we compare the constitution of a neutral salt with that of the hydrated acid by which it is formed, we find the positive result to be the substitution of a metal for the hydrogen of the latter, thus, SO' + HO gives with zinc SO' + ZnO; and where a metal is acted on by an hydrated acid, the hydrogen is thus evolved either directly as gas, or it reacts on the elements of the acid and gives rise to secondary products which are evolved, such as sulphurous acid, nitric oxide, &c. In all cases we may consider the action of a metal on a hydrated acid, to be primarily the elimination of hydrogen and the formation of a neutral salt. But in this respect the action becomes completely analogous to that of the metal on a hydracid, except that in the latter case a haloid salt is formed, and hence we assimilate the two classes in

constitution by a very simple arrangement of their formulæ.

There are, however, a number of acids which may be obtained in a dry and isolated form, as the sulphuric, the silicic, the telluric, the stannic, the arsenic, the phosphoric, &c., and when they combine with bases, it is most natural to consider the union as being direct, and that the salt contains acid and base really as such. This is accordingly the strongest point of the ordinary theory. But other and important circumstances intervene. These acids, although they may be obtained free from water, yet in that state they combine with bases but very feebly, and require a high temperature in order to bring their affinities into play. On the other hand, in all cases where these bodies manifest their acid characters in the highest degree, they are combined with water, as in oil of vitriol and phosphoric acid, and when expelled from combination with a base, they immediately enter into combination with water in an equivalent proportion. Thus where phosphate of lime is decomposed by oil of vitriol, it is not phosphoric acid (POs) which is found in the liquor, but its



ric acid in vapour, and yet cold barytes and oil of a tensity as to produce ignition, whilst the barytes m combine with the dry sulphuric acid. The water, the festation of strong acid properties, and it does not a merely as a base. What, then, is the constitution of the strong acid properties.

When muriatic acid (H.Cl.) acts on zinc, the mand hydrogen is expelled, and if, in place of zinc, c is the same, except that the hydrogen combining w water; HCl and ZnO giving ZnCl and HO. Now menta SO'H combined together, when put in cont SO'Zn is formed, and with ZnO and SO'H, there is set free. In both cases, of which the former may haloid salts and the latter of all salts formed by ox ment which is removable by a metal, precisely as a ther, as indeed from the real metallic character of occur in this case. Every acid may, therefore, be a combined with an electro-negative element, which is fluoring, or may be compound, as cyanogen, NC2, or as occurs in the great inspority of cases, its eleminant together when in combination. Thus oil of a HO, but consists of hydrogen united to a compound does not contain NO' and HO, but consists of hydreal NO's, and the acetic acid is written C'H'O' + 1

The elegance and simplicity with which the ladeduced from these principles is really remarkable a fact substantiated by experiment, that in neutral of acid were proportional to the number of equival the ordinary theory gave no indication of why this sarily from the principles of the newer theory. The an acid, M denoting the metal of the oxide, and R is ing action is

M + O and H + R, produce H +

and in the neutral salt there is an equivalent of eac oxide, in order that water shall be formed, and so the reaction is that

 $M^{4} + O^{3}$  and 3 (H + R), produce 3 (H

a sesqui-compound being formed perfectly analogous ber of atoms of acid, 3 (H + R), is equal to the nus base (M<sup>2</sup>O<sup>3</sup>), because that number of atoms of hydro position of the base. In like manner for a deutoxid ence there is no other proof than their utility in supporting this view, becomes more powerful as an objection, when we proceed to apply its principles to the salts of phosphoric acid. For it has been already described, that this acid forms three distinct classes of salts, all neutral, and which have their origin in the three hydrated states of the phosphoric acid. These states are written on the two views as follows:—

|                 | Old Theory.   | New Theory.  |
|-----------------|---------------|--------------|
| Monobasic acid, | $PO^{5} + HO$ | $PO^6 + H$   |
| Bibasic acid,   | PO' + 2HO     | $PO^7 + H^2$ |
| Tribasic acid.  | $PO^5 + 3HO$  | $PO_8 + H_3$ |

Now it appears very useless, where the older view accounts so simply for the properties and constitution of these salts, to adopt so violent an idea, as that there are three distinct compounds of phosphorus and oxygen which no chemist has ever been able to detect. But here again other circumstances must be studied; first, the difference of properties of phosphoric acid, in its three states, is totally inexplicable, on the idea of their being merely three degrees of hydration. Nitric acid forms three hydrates, but when neutralized by potash, it always gives the same saltpetre; sulphuric acid forms two perfectly definite hydrates, but with soda forms always the same glauber's salt; whilst phosphoric acid, when neutralized by soda, gives a different kind of salt according to the state it may be in. Also, the permanence of these conditions of phosphoric acid is a powerful proof that they do not consist in the adhesion of mere water. The idea that the phosphoric acid is a different hydracid in each of its three conditions, on the other hand, not merely explains the fact of these differences of properties, but it renders the formation of bibasic and tribasic salts, which is such an anomaly on the old theory, a necessary consequence of the new, for the phosphoric salt radicals, PO<sup>6</sup>, PO<sup>7</sup>, and PO<sup>8</sup>, differ not merely in the quantity of oxygen they contain, but are combined with different quantities of hydrogen, and hence in acting on metallic oxides (bases), there is a different number of atoms required for each to replace the hydrogen and form water. Thus—

PO6.H and NaO give HO and PO6.Na. monobasic phosphate of soda, PO7.H2 and 2NaO give 2HO and PO7.Na2. bibasic phosphate, PO6.H3 and 3NaO give 3HO and PO8.Na3. tribasic phosphate.

A circumstance which gives additional reason to infer that the water is not merely as base in the phosphoric acid, is the following: if it were so, then it should be most completely expelled by the strongest bases, and the bibasic and tribasic phosphates of the alkalies should be those least likely to retain any portion of the basic water; but the reverse is the fact; whilst oxide of silver, a very weak base, is that which most easily and totally replaces the water. On the idea, however, of hydracids, this is easily understood, for the oxide of silver is one most easily reduced by hydrogen, and consequently one on which the action of a hydrogen acid, as  $PO^4 + H^3$ , or  $PO^7$ 

+ H<sup>2</sup>, would be most completely exercised.

A remarkable verification of this theory has been recently found in the decomposition of solutions of the oxysalts in water, by voltaic electricity. It has been already explained (pp. 314 et seq.), that it requires the same quantity of electricity to decompose an equivalent of any binary compound, such as iodide of lead, chloride of silver, muriatic acid, or water. Now, if we dissolve sulphate of soda in water, and pass a current of voltaic electricity through that solution, we have water decomposed, and also the glauber's salt; oxygen and sulphuric acid being evolved at one pole, and soda and hydrogen at the other. Here, on the old view, the electricity performs two decomposing actions at the same time, and, as it thus divides itself, its action on each must be lessened, and the quantity of each decomposed be diminished, so that the sum should represent the proper energy of the current. On measuring these quantities, however, the result is totally different, the quantity of sulphate of soda decomposed is found to be equal to the full duty of the current, and an equivalent of water appears to be decomposed in addition. It is quite unphilosophic to imagine, that the strength of a current should be thus suddenly doubled, and a simple and sufficient explanation of it is found in the new theory of salts. The sulphate of soda in solution having the formula Na.SO4 is resolved by the current into its elements. Na and SO4, as chloride of sodium would also be; the sodium, on emerging at the negative electrode, from the influence of the current, instantly decomposes water, and soda and hydrogen, of each an equivalent, are evolved; at the positive electrode the compound radical SO4 also decomposes water, and produces H.SO4 and O. The appearance of the oxygen and hydrogen is thus but secondary, and the body really decomposed by the current is only Na SO4.

In the case of the salts of such metals as do not decompose water, the phenomena

are much more simple. Thus a solution of sulphate of copper, when decommely the battery, yields metallic copper at the negative, and sulphure and are can at the positive electrode, and the quantity of copper separated represent the energy of the current which has passed, for the salt being Cu SO', it and the into its elements, but SO4 reacting on the water, produces H SO4 and 0 u 20 ... tive electrode. On the old view, it was supposed that water and sulptain f were both decomposed, oxygen and acid being evolved at one are and no en per and hydrogen being separated at the other, which reacting professions the metal. Such an explanation, however, is directly opposed to the awd to in nite action of electricity, and cannot be received

In the case of sclutions of chlorides or iodides, where there can be no destrict relations of the elements, the results of voltage decomposition are precise; Chloride of copper gives simply chlorine and copper, no water being descent. Chloride of sodium or iodide of potassium give chlorine or iodine at the controle, and alkali and hydrogen at the other, the evolution of three last being rest

by the action of the metallic basis on the water of the solution

Professor Daniell, to whom these important electro-chemical researche in the

considers the truth of the binary theory of salts to be fully established in their lift this theory be adopted, a profound change in our nomenclature of salt will come necessary. Graham has proposed that the mame of the sait races at city formed by prefixing to the word oxygen the first word of the ordinary rame # ... class of salts, and that the salts be termed by changing orugen into anite the SO4. sulphatoxygen, gives sulphatoxides, the sulphates. NO4 n tratexygen mitratexides, the nitrates, and so on; but I consider that the form of some states. proposed by Dantell deserves the preference. It has been described possible for a large formations; acting on this, Daniell proposes to term the electro-negative element of sulphates oxysulphion, that of the nitrates oxysulphion, and so on, and the minutes of sulphates oxysulphion of supposes to term the electro-negative element of sulphates oxysulphion, that of the nitrates oxysulphion, and so on, and the minutes of supposes of suppo termed expectation of copper, expectation of sodium, &c. It would be describe. however, for a long time, to introduce these names only where theoretical rations rendered their employment decidedly useful, and hence, in all fature decidedly tion of the salts, I shall make use of the language of our ordinary views and multitheir preparation and composition without any reference to the discussion a wini

we have been engaged

The general adoption of the binary theory of salts has deprived of much of man terest and importance a question, which some years since was very ingree at the cussed, viz -whether, in the formation of double salts, the salts which man be to same relation to each other that acid and base were then thought to have The was supposed that the electro-negative qualities of aulphuric acid beat comtrolled by oxide of copper than by potash, the aikaline suiphate acted as a tem be the sulphate of copper, when these two salts combined to form the double of potash and copper, and so on in other instances; but in addition to the constitution of the salts militates are wellview, we have the positive evidence that, first, these double saits are formed that combination merely, but by replacement of the constitutional water of the same of the copper or magnesian class, which water nobody would contend to act a 22 as a base; and second, that when a solution of such a double salt is dec moves the battery, the two salts are not separated as if they were acid and base to 1" decomposed independently in the proportions of an equivalent of each, manage-

gether the sum of the chemical energy of the current.

A similar idea was advocated by Bonsdorff regarding the double chloride is the complete that it &c. He proposed to consider the chlorides of gold, platina, mercury &c as t rine acids, and those of potassium, &c , as chlorine bases, and so with the This view, however, although at first very extensively adopted, has given at the gradual growth of knowledge. There is no analogy between a dry 172/201 and a chloride, but the chlorides are in perfect analogy with the scutt. As Thus CuCl does not resemble SO3, but Cu.SO4 and CuCl + KCl 18 and 5.44 to SO3 KO, but to the double salt CuSO4 + KSO4. Bonsdorff's idea was to counter to the direction of truth, he sought to bring all salts under the one be extending to all the constitution of exygen acids and exygen bases, which  $z_{r}$   $z_{r}$ gress of science has led us to the opposite generalization of reducing all said ' 2'

simple haloid type."

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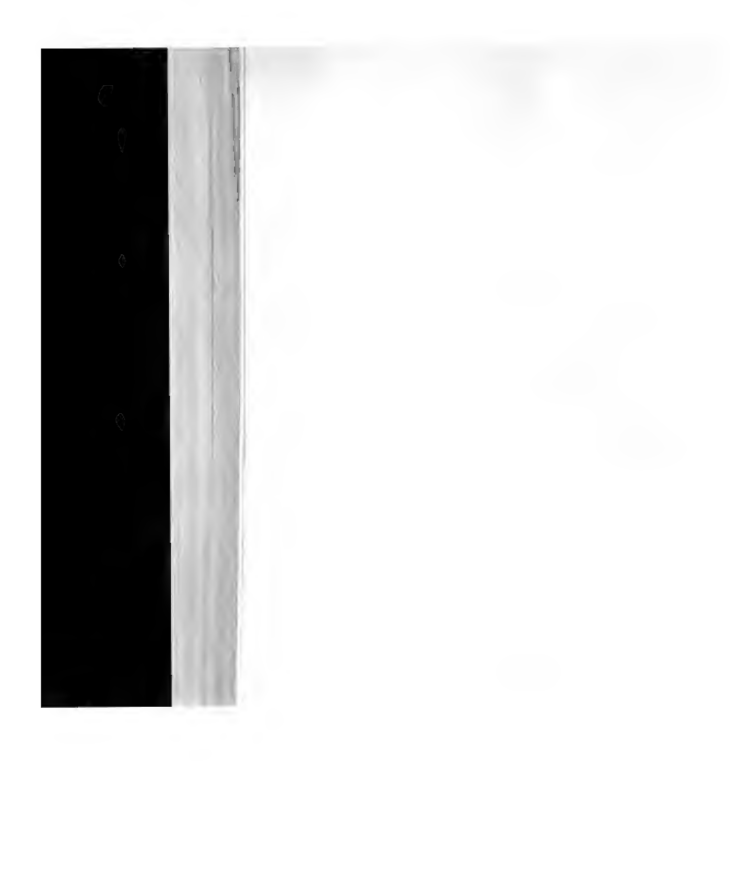
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